Enhancer Priming Enables Fast and Sustained Transcriptional Responses to Notch Signaling.

Permalink
https://escholarship.org/uc/item/0hm7g7ws

Journal
Developmental cell, 50(4)

ISSN
1534-5807

Authors
Falo-Sanjuan, Julia
Lammers, Nicholas C
Garcia, Hernan G
et al.

Publication Date
2019-08-01

DOI
10.1016/j.devcel.2019.07.002

Peer reviewed
Enhancer Priming Enables Fast and Sustained Transcriptional Responses to Notch Signaling

Highlights

- Notch-responsive enhancers are activated synchronously in Drosophila embryos
- Notch levels modulate transcription by changing burst size
- Notch elicits stochastic versus deterministic activation depending on cell context
- Priming by Twist and Dorsal promotes fast and sustained response to Notch

In Brief

Falo-Sanjuan et al. use live imaging to investigate how enhancers detect and respond to Notch signaling in real time in vivo. They discover that responses are synchronized and sustained in the normal context, due to local transcription factor priming, and that elevated Notch activity increases transcription burst size rather than frequency.

Authors

Julia Falo-Sanjuan, Nicholas C. Lammers, Hernan G. Garcia, Sarah J. Bray

Correspondence

sjb32@cam.ac.uk
Enhancer Priming Enables Fast and Sustained Transcriptional Responses to Notch Signaling

Julia Falo-Sanjuan,1 Nicholas C. Lammers,2 Hernan G. Garcia2,3,4,5 and Sarah J. Bray1,6,*

1Department of Physiology, Development and Neuroscience, University of Cambridge, Downing Street, Cambridge CB2 3DY, UK
2Department of Physics, UC Berkeley, Berkeley, CA 94720, USA
3Department of Physics, UC Berkeley, Berkeley, CA 94720, USA
4Department of Molecular and Cell Biology, UC Berkeley, Berkeley, CA 94720, USA
5Institute for Quantitative Biosciences-QB3, UC Berkeley, Berkeley, CA 94720, USA
6Lead Contact
*Correspondence: sjb32@cam.ac.uk
https://doi.org/10.1016/j.devcel.2019.07.002

SUMMARY

Information from developmental signaling pathways must be accurately decoded to generate transcriptional outcomes. In the case of Notch, the intracellular domain (NICD) transduces the signal directly to the nucleus. How enhancers decipher NICD in the real time of developmental decisions is not known. Using the MS2-MCP system to visualize nascent transcripts in single cells in Drosophila embryos, we reveal how two target enhancers read Notch activity to produce synchronized and sustained profiles of transcription. By manipulating the levels of NICD and altering specific motifs within the enhancers, we uncover two key principles. First, increased NICD levels alter transcription by increasing duration rather than frequency of transcriptional bursts. Second, priming of enhancers by tissue-specific transcription factors is required for NICD to confer synchronized and sustained activity; in their absence, transcription is stochastic and bursty. The dynamic response of an individual enhancer to NICD thus differs depending on the cellular context.

INTRODUCTION

Genes respond to external and internal cues through the actions of transcription factors and effectors of signaling pathways. Gene regulatory regions, termed enhancers, integrate information from these inputs to produce an appropriate transcriptional output. During development, decisions may occur in a matter of minutes, but as the transcription dynamics have rarely been analyzed in vivo in real time, we know little about how recipient enhancers decipher the signals. For example, enhancers could respond in a digital manner, working as simple on-off switches, or as analog devices, operating as a rheostat so that signal levels can modulate the output (Blackwood and Kadonaga, 1998; Garcia et al., 2013; Lammers et al., 2018). In either case, they must also have the capability to detect and transduce key parameters to the transcription machinery, such as signal duration and thresholds.

With the advent of precise and quantitative methods to measure transcription, such as single molecule fluorescence in situ hybridization (smFISH) or live imaging, it has become evident that transcription is not a continuous process. Instead, transcribing genes undergo bursts of initiation that are often separated by inactive intervals (Chubb et al., 2006; Golding et al., 2005). Bursting is thought to occur because dynamic enhancer-promoter activation leads to episodic polymerase release. One consequence of this is that factors modulating the levels of transcription can do so by changing either the frequency with which a burst occurs (measured by the gap between bursts) or the size of each burst (measured by changes in burst duration and/or amplitude). To date, bursting frequency rather than burst duration or amplitudes seems to be the major parameter modulated in different species and contexts (So et al., 2011; Senecal et al., 2014; Xu et al., 2015; Desponts et al., 2016; Padovan-Merhar et al., 2015; Lammers et al., 2018; Berrocal et al., 2018). For example, enhancers controlling early patterning genes in Drosophila embryos produce similar bursting size but have different bursting frequencies, which can be attenuated by the presence of insulators (Fukaya et al., 2016). Similarly, steroids increase the bursting frequency of target enhancers (Larson et al., 2013; Fritzsch et al., 2018). However, it remains to be discovered whether all transcription factors alter transcription dynamics in this way and specifically whether it is these or other properties that are modulated by developmental signals to confer appropriate outputs in the in vivo setting of a developing organism.

Transcriptional bursting is thought to make an important contribution to heterogeneity in transcriptional activity between cells (Raj and van Oudenaarden, 2008). For example, in cells exposed to estrogen, response times for transcription activation were highly variable with no coherent cycling between active and inactive states (Fritzsch et al., 2018). Stochastic transcriptional behavior is also of key importance for differentiation of photoreceptors in Drosophila eyes (Wernet et al., 2006), of hematopoietic cells in mice (Chang et al., 2008; Ng et al., 2018), and of neuronal cells in zebrafish retina (Boije et al., 2015), however,
Figure 1. Synchronous Activity of Two Notch-Responsive Enhancers
(A) Diagrams illustrating the MS2 strategy for live imaging of transcription (top) and the location of mesectoderm (MSE) and neuroectoderm (NE) enhancers in E(spl)-C (m5/m8; m8NE) and single minded (sim) (bottom). Arrows indicate transcription start-sites, boxes in lower panel indicate promoters (white), non-coding (light gray) and coding (dark gray) transcribed regions.
(B) Diagram of a blastoderm Drosophila embryo, indicating mesodermal Delta expression (pink), which activates Notch in flanking cells (green dots) to specify the MSE. Image: transcription from m5/m8 detected by MCP-GFP accumulation in bright puncta (green), nuclei are labeled by His2Av-RFP (blue).
(C) Tracked expression from m5/m8 and sim reporters. Top panels: tracked nuclei false-colored by total signal levels, proportional to total mRNA production.
Bottom panels: single frames with tracked nuclei shaded according to maximum pixel intensity.
(D) Profiles of m5/m8 and sim fluorescence from individual nuclei that exhibit “sustained” activity.
(E) Heatmaps representing fluorescence profiles of m5/m8 and sim in all MSE nuclei during nc14 (scale as indicated with blue, no expression; yellow, high expression; black indicates periods where nuclei were not tracked).
(F) Distributions of onsets and end points of transcription from m5/m8 and sim in the MSE.
(G) m5/m8 and sim produce similar average temporal profiles. Mean fluorescent intensity of MCP-GFP puncta at indicated times in nc14.
while it is an attractive feature for promoting heterogeneity, inherent transcriptional variability could be extremely disruptive in developmental processes where a coordinated response of many cells is required to pattern specific structures. In some cases, this may be circumvented by averaging mechanisms that allow cells to produce homogeneous patterns of gene expression (Little et al., 2013) that include mRNA diffusion in Drosophila syncytial embryos (Bothma et al., 2018). However, it is only in rare circumstances that mRNA diffusion can operate, and it is unclear whether other averaging mechanisms would be effective over shorter time intervals. To effectively achieve reproducible patterns, cells must therefore overcome the variability that is inherent in transcriptional bursting and stochastic enhancer activation.

Notch signaling is a highly conserved developmental signaling pathway that is deployed in multiple contexts. It has the unusual feature that the Notch intracellular domain (NICD) transduces the signal directly to the nucleus, when it is released by a series of proteolytic cleavages precipitated by interactions with the ligands. NICD then stimulates transcription by forming a complex with the DNA binding protein CSL and the co-activator Mastermind (Mam) (Bray, 2006). The lack of intermediate signaling steps and amplification makes this a powerful system to investigate how signals are deciphered by responding enhancers. Furthermore, there may be differences in the levels and dynamics of NICD produced by different ligands (Nandagopal et al., 2018). However, although its role as a transcriptional activator is well established, at present, we know little about how enhancers respond to NICD in the real time of developmental decisions. For example, do enhancers operate as simple switches, detecting when NICD crosses a threshold, or are they sensitive to different levels of NICD, in which case does NICD, like other factors, modulate bursting frequency? Nor do we know what sequence features in the responding enhancers confer the output properties, although enhancers with paired CSL motifs (referred to as SPS motifs) (Bailey and Posakony, 1995; Nam et al., 2007), whose precise spacing could favor NICD dimerization, are suggested to yield the strongest responses (Nam et al., 2007).

In order to determine how enhancers respond to Notch activity in real time, we have used the MS2-MCP system to visualize nascent transcripts in Drosophila embryos. To do so, we used two well-characterized Notch-responsive enhancers that drive expression in a stripe of mesoderm (MSE) cells and analyzed their transcription profile over time at the single cell level. Strikingly, all MSE cells initiated transcription within a few minutes of one another, and once active, each nucleus produced a sustained profile of transcription. By manipulating NICD levels and altering key motifs within the enhancers, we uncover two key principles. First, the ability of NICD to confer synchronized and sustained activity in MSE requires that the enhancers be primed by tissue-specific transcription factors. In their absence, MSE enhancers confer stochastic bursty transcription profiles, demonstrating that different response profiles can be generated from a single enhancer according to which other factors are present. Second, changing Notch levels modulate the transcription burst size but not the inter-burst periods, in contrast to most current examples of enhancer activation. These two key concepts are likely to be of general importance for gene regulation by other signaling pathways in developmental and disease contexts.

RESULTS

Synchronized and Sustained Enhancer Activation in Response to Notch

To investigate how Notch signals are read out by an enhancer in real time, we focused on well-characterized MSE enhancers from the Enhancer of split-Complex (E(spl)-C) (known as m5/m8) and from singleminded (sim) (Morel and Schweisguth, 2000; Cowden and Levine, 2002; Zinzen et al., 2006a). These direct expression in two stripes of MSE cells during nuclear cycle 14 (nc14) when Notch is activated in response to Delta signals from the presumptive mesoderm (ME) (Figures 1A and 1B) (Morel et al., 2003; De Renzis et al., 2006; Zinzen et al., 2006a). The MSE converges to the midline during gastrulation, ultimately forming CNS midline precursors similar to the vertebrate floorplate. To visualize transcription from MSE enhancers in real time and define the response properties conferred by a defined enhancer DNA sequence, they were inserted into MS2 reporter constructs comprising the even-skipped promoter (peve), 24 MS2 loops, and lacZ (Figure 1A). When combined with MCP-GFP in the same embryos, nascent transcription from the MS2 reporters was detected by the accumulation of MCP-GFP in nuclear puncta, whose fluorescence is directly proportional to the number of transcribing mRNAs at any time point (Figures 1A and 1B) (Garcia et al., 2013). In this way, levels of transcription can be followed over time in each cell by tracking the puncta relative to nuclei.

Visualizing transcription in real time revealed that m5/m8 and sim were both activated in all MSE cells within a narrow time window (~10 min) in nc14 (Figures 1C, 1E, and 1F; Videos S1 and S2). Activity was then maintained in these nuclei throughout the remaining period of nc14 as embryos underwent gastrulation. Both m5/m8 and sim exhibited what we refer to as “sustained activity” because each punctum retained high levels of fluorescence rather than exhibiting clearly distinct bursts (Figure 1D), although we note that the resolution of bursting events is limited by the time each polymerase takes to complete transcription (estimated as 1.6–2.5 min for these reporters) (Fukaya et al., 2017). Transcription then ceased after 30–50 min, with less synchrony than at the onset (Figure 1F). Identical response profiles were obtained when the m5/m8 reporter was inserted at a different genetic locus (Figures S1A and S1B).

Sustained activity is a feature of m5/m8 and sim and not a general property of Notch-responsive enhancers at this stage, as a neuroectodermal enhancer from E(spl)m8-bHLH (m8NE, Figure 1A) produced profiles where individual bursts of activity

(‡) Transcription from m5/m8 is curtailed in embryos lacking zygotic Delta (Dl) and abolished in embryos lacking neuralized (neur).

Gray trace, m5/m8 profile in wild-type embryos from (G). In (G) and (H), mean and SEM of all MSE cells are shown. n = 3 (m5/m8), 3 (sim), 2 yn5/m8; Dl), 2 yn5/m8; neur) embryos. In this and other figures, the peve promoter was used in all reporters unless otherwise specified. See also Figure S1 and Videos S1 and S2.
were clearly resolved, which we refer to as “bursty” (Figures S1A–S1C). Furthermore, even though profiles produced by m5/m8 and sim were continuous, their amplitude fluctuated, likely reflecting episodic polymerase release. Overall, however, the m5/m8 and sim response profiles were highly coordinated temporally (Figures 1E and 1F). Indeed, the mean profile of all MSE cells analyzed was almost identical for the two enhancers (Figure 1G). This is remarkable given that they contain different configurations of binding motifs and implies that MSE cells undergo a highly synchronized period and level of Notch signaling.

To assess the relative contributions of the enhancer and promoter to response profiles, we next tested consequences of substituting different promoters with m5/m8 and sim, inserting the reporters at the same genomic position to ensure comparability. First, when peve was replaced by a promoter from sim (psimE), both m5/m8 and sim produced lower levels of transcription, but their overall temporal profiles remained similar (Figure S1D). Second, when we combined m5/m8 with another heterologous promoter, hsp70, or with four promoters from the E(spl)-C locus, mean levels of transcription were again affected without changing the overall temporal profile or expression pattern (Figure S1E). Notably, even in combinations yielding lower levels, e.g., pm6 (Figure S1E), the transcription profiles remained sustained rather than breaking down into discrete bursts (Figure S1F). Although the results suggest there could be an underlying enhancer-promoter compatibility at the sequence level (Figure S1E) (Zabidi et al., 2015), there was no obvious relationship between the mean levels of transcription produced by a promoter and the presence or absence of sequence motifs for factors associated with promoter accessibility, such as Zelda or Trithorax-like (Blythe and Wieschaus, 2016). Nor was there a correlation between promoter activity with m5/m8 in the MSE and that with a heterologous developmental enhancer in Drosophila S2 cells (Arnold et al., 2016). However, since the promoter substitutions had no effect on temporal profiles, it argues that the enhancers are the primary detectors of Notch activity.

To verify that MSE transcription was Notch dependent, we measured transcription from m5/m8 in embryos where Notch activity was disrupted by mutations. Embryos lacking Neuralized, an E3 ubiquitin ligase required for Delta endocytosis that is critical for Notch signaling (Morel et al., 2003; De Renzis et al., 2006), had no detectable transcription from m5/m8 in the MSE (Figure 1H). Likewise, m5/m8 activity was severely compromised in embryos carrying mutations in Delta. Because Delta protein is deposited in the egg maternally (Kopczynski et al., 1988), these embryos contained some residual Delta, which was sufficient for a few scattered MSE cells to initiate transcription (Figure S1G). However, their transcription ceased prematurely, within <20 min (Figures 1H and S1G). Together, these results confirm that the enhancers require Notch signaling for their activity in the MSE, in agreement with previous studies (Morel and Schweisguth, 2000; Zinzen et al., 2006a), and further show that continued Notch signaling is needed to maintain transcription, arguing that the MSE enhancers also detect persistence of NICD.

**Coordinated Activity of Enhancers within Each Nucleus**

Although m5/m8 and sim confer well-coordinated temporal transcription profiles, their precise time of activation shows some cell-to-cell variability (Figure 1F). To investigate whether this variability reflects stochastic variations in transcription (intrinsic variability) or differences in signaling from Notch (extrinsic variability) (Elowitz, 2002; Raser and Shea, 2006), we monitored expression from two identical alleles of the MS2 reporters (Figure 2A). Transcription from these two physically unlinked loci was detected as distinct puncta in each nucleus, which could be tracked independently. We found a remarkable synchrony in the onset of transcription from both alleles of a given enhancer (Figure 2B). More than 80% of cells initiated transcription from both alleles within 5 min (Figure S2C). This contributes to ~6%–30% of the total variability (Figure 2D), indicating that most onset variability was due to extrinsic factors. Transcription was extinguished less synchronously (Figures 2B and S2A), but this intrinsic variability was still much less than that between cells (Figure 2D).

Although two alleles in the same cell gave overall similar profiles, their fine-grained spikes and troughs were not synchronized (Figure 2A), as expected if transcription from two different loci is largely uncorrelated (Harper et al., 2011; Little et al., 2013; Fritzsch et al., 2018). However, their fluorescence intensities displayed a small but significant positive correlation (R² ~ 0.35) (Figure S2B). This argues that the enhancers at the two alleles operate independently while being coordinated by the same extrinsic information. Strikingly, when m5/m8 and sim were present in trans in the same cell, there was also comparatively little variation in their onset times (Figures 2C, 2D, and S2A). Thus, the properties of m5/m8 and sim ensure that they reliably detect extrinsic information in the form of Notch activity, initiated within a 5- to 10-min time window so that their activation is remarkably synchronized within each nucleus.

**Enhancers Detect Signal Thresholds and Context**

The m5/m8 and sim enhancers appear to act as “persistence detectors,” driving transcription as long as Notch signal(s) are present. They may simply detect when Notch reaches a threshold (digital encoding) or they could be sensitive to Notch activity levels (analog encoding). To distinguish these possibilities, we supplied ectopic NICD using the stripe 2 regulatory enhancer from even-skipped (eve2-NICD). This produces an ectopic stripe of NICD, orthogonal to the MSE (Figure 3A) (Kosman and Small, 1997; Cowden and Levine, 2002), which was sufficient to produce ectopic expression from both m5/m8 and sim (Videos S3 and S4).

Whereas expression from m5/m8 and sim was almost identical in wild-type embryos, clear differences were revealed by ectopic NICD. First, m5/m8 was activated throughout the dorsal eve2 domain whereas sim only responded in a 3- to 4-cell region close to the MSE (Figure 3B), consistent with previous observations (Cowden and Levine, 2002; Zinzen et al., 2006a). Second, although both enhancers initiated transcription prematurely because the ectopic NICD was produced from early nc14 (Bothma et al., 2014), m5/m8 was switched on significantly earlier than sim (Figures 3G and 3H). Given that both enhancers are exposed to the same temporal pattern of NICD, this difference in initiation times implies that they respond to different thresholds of NICD. Therefore, we hypothesize that m5/m8 and sim respond at the same time in wild-type embryos because the normal ligand-induced signaling leads to a sharp increase in NICD.
We also detected differences in the response dynamics of m5/m8 according to location. Nuclei close to the MSE stripe (in the neuroectoderm, NE) exhibited strong activity, with a temporal pattern resembling that in the MSE (Figure 3C, bottom). In contrast, nuclei in more dorsal regions (dorsal ectoderm, DE) underwent resolved bursts of transcriptional activity (Figure 3C, top). Ectopic NICD also induced “bursty” expression from sim in the ME but was not capable of turning on m5/m8 in that region (Video S5).

“Bursty” m5/m8 transcription in the DE was also associated with more stochastic activation. In embryos with two m5/m8 alleles, both were activated in response to eve2-NICD in most MSE and NE nuclei, whereas only a single allele was active at any one time in most DE nuclei (Figures 3D and 3F; Video S6). Furthermore, in the few DE nuclei where both alleles became active, there was greater variability in onset times and the profiles were less coordinated (Figures 3E and S2D). The positional differences in dynamics suggest that intrinsic cellular conditions, likely expression levels of specific transcription factors, influence the way that enhancers “read” the presence of NICD. Such factors must therefore have the capability to modulate the dynamics of transcription.

**Notch Activity Tunes Transcription Burst Size**

To further test how Notch-responsive enhancers respond to doses of signal, we introduced a second eve2-NICD transgene. MSE transcription from sim in 2xeve2-NICD embryos initiated earlier and achieved higher levels than with 1xeve2-NICD (Figure 4A, left). This agrees with the hypothesis that sim responds to higher thresholds of NICD, as nuclei will reach a given concentration of signal more quickly in embryos with 2xeve2-NICD. The mean levels of transcription increased in ME as well as in MSE regions (Figures 4A–4C), further indicating a dose-sensitive response. In contrast, levels and onset of MSE transcription from m5/m8 did not significantly change in 2xeve2-NICD embryos (Figure 4A, right). This saturation in output from m5/m8 only occurred in the MSE, as the more stochastic activity in the DE remained sensitive to increases in NICD, being detected in a greater proportion of cells and over longer periods (Figure S4A).
Figure 3. Effects of Ectopic NICD on Temporal Transcription Profiles Reveal that Enhancers Have Different Thresholds

(A) Strategy for producing ectopic NICD using eve2, with schematics depicting expression (purple shading) relative to MSE (green) and DV regions where effects on transcription were quantified.

(legend continued on next page)
To distinguish different models for how NICD confers a dose-sensitive response, we took two strategies to analyze its effect on transcriptional bursting dynamics and focused on regions where individual bursts of transcription were resolved. Both approaches assume a two-state model where the enhancer is switched between an OFF and ON state with switching rates $K_{on}$ and $K_{off}$ and confers transcription initiation rate $r$ in the ON state (Figure 4E) (Peccoud and Ycart, 1995; Larson et al., 2009). In the first approach, we directly measured bursting amplitude, off period between bursts and bursting length as approximations for $r$, $K_{on}$, and $K_{off}$, respectively (Figure 4E). In most previous enhancers analyzed in this way, the off period is the most affected, leading to changes in the bursting frequency (Peccoud et al., 2016; Fritsch et al., 2018; Lammers et al., 2018). However, when we quantified the effect from different doses of NICD on $sim$ in the ME, we found that bursting length consistently increased with higher amounts of NICD, whereas off periods between bursts remained constant (Figures 4D and 4F). This indicates that the main effect of NICD is to keep the enhancer in the ON state for longer—i.e., decreasing $K_{off}$—rather than increasing the frequency with which it becomes active—i.e., increasing $K_{on}$. The bursting amplitude also increased with $1x$eve2-NICD but this was not further enhanced by $2x$eve2-NICD (Figures 4D and 4F). Overall, therefore, increasing levels of NICD in the ME result in $sim$ producing an increase in transcription burst size (duration $\times$ amplitude) rather than an increase in the frequency of bursts. A similar increase in burst size in response to the dose of NICD (Figures S4A–S4C) occurred with other regions and enhancers ($m5/m8$ DE and $m8$NE ME), suggesting that it is a general property of these Notch-responsive enhancers.

We developed a second approach, based on the noise properties of transcription, to analyze the changes in dynamics even where single bursts of activity could not be defined. To do so, we used a mathematical model of transcription to account for the initiating mRNA molecules (Figure S3A). Using derivations from the mathematical model and testing them in simulations, we looked for signatures that would be produced if the mean of initiating mRNAs (equivalent to the mean fluorescence from $m5$ puncta) were increasing due to changes in $r$, $K_{on}$, or $K_{off}$. This showed that the effects on the Fano factor ratio between the two conditions and on their autocorrelation function (ACF) could be used to correctly predict which of the parameters could account for the increase (Figure S3B; STAR Methods). First, we tested the modeling approach with the data from the promoter swap experiments. Analyzing the differences in the mean indicated that they are most likely due to increases in $r$ (Figure S4D), as expected if promoters influence the rate of polymerase release but not enhancer activation per se. When we then applied the model to data from the transcription profiles produced by different doses of NICD in the ME, results were most compatible with the causal effect being an increase in $r$ or a decrease in $K_{off}$ (Figure S4E) depending on which two conditions were compared. Thus, this second approach also indicated that NICD elicits an increase in burst size rather than in burst frequency. Both approaches therefore converge on the model that above the critical threshold level of NICD, further increases in NICD levels prolong the period that each enhancer remains in the ON state.

Finally, we used an enhancer-promoter combination that produced higher mean levels ($m5/m8$-pm5, Figure S1E) to investigate whether the saturation that occurred with ectopic NICD was due to the $peve$ promoter having achieved maximal initiation rate. Strikingly, the substitution of pm5 for peve did not result in significantly higher maximal levels in the presence of eve2-NICD (Figure S4F), although it did in wild-type embryos (Figure S1E). This indicates that the saturation of the response with higher levels of NICD stems from the $m5/m8$ enhancer rather than the promoter and argues that enhancers reach a maximal “ON” state that they cannot exceed even if more NICD is provided.

**Paired CSL Motifs Augment Burst Size, Not Threshold Detection**

$m5/m8$ and $sim$ enhancers both respond to NICD but they initiate transcription at different thresholds. How is this encoded in their DNA sequence? A prominent difference is that $m5/m8$ contains a paired CSL motif (so-called SPS motif) whereas sim does not (Figure S5A). To test their role, we replaced two CSL motifs in $sim$ with the SPS motif from $m5/m8$ and conversely perturbed the SPS in $m5/m8$ by increasing the spacing between the two CSL motifs (Figure S5A). As SPS motifs permit cooperative binding between two NICD complexes, we expected that enhancers containing an SPS motif ($sim^{SPS}$ and $m5/m8^{SPS}$) would exhibit earlier onsets of activity than their cognates without (sim and $m5/m8^{SSPS}$). However, this was not the case for either $sim$ and $sim^{SPS}$ (Figures 5A and 5B) or $m5/m8$ and $m5/m8^{SSPS}$ in either wild-type or eve2-NICD embryos (Figures S5D and S5E). These profiles suggest that the SPS motifs are not responsible for the difference in the threshold levels of NICD required for $m5/m8$ and $sim$ activation.

Changes to the CSL motifs did, however, affect mean levels of activity. $sim_{SPS}$ directed higher mean levels of activity compared to $sim$ in both wild-type and eve-NICD embryos (Figures 5A and S5B). Conversely, $m5/m8^{SSPS}$ directed lower levels compared to $m5/m8$ (Figure S5D). Analyzing the traces from $sim$ in the ME, where cells undergo resolved bursts of transcription,
**Figure 4. Notch Produces a Dose-Sensitive Response by Regulating Transcription Burst Size**

(A) Mean levels of transcription from sim (left) and m5/m8 (right) in the MSE with (2x eve2-NICD), compared to 1x eve2-NICD and to the wild type.

(B) Heatmaps depicting sim activity in ME nuclei in three conditions as indicated. Note the different scale range compared to Figure 3G.

(C) Mean levels of transcription from sim in ME produced by different doses of NICD.

(D) Examples of transcription traces from single ME nuclei in the wild type, 1x eve2-NICD, and 2x eve2-NICD. Burst periods are marked with a gray line.

(E) Schematic of the model: an enhancer cycles between ON and OFF states and produces mRNA when ON. Changes in bursting amplitude, off period, and bursting duration correlate with changes in kinetic constants $r$, $K_{on}$, and $K_{off}$.

(legend continued on next page)
revealed that the SPS motif (simSPS) led to larger burst sizes—i.e., increased the amplitude and the duration—compared to wild-type sim (Figures 5C and 5D). Conversely, the continuous profile produced by m5/m8 in the MSE was broken into smaller bursts when the SPS was disrupted (Figures S5F and S5G). The effects on bursting size are similar to those seen when the dose of NICD was altered, suggesting that enhancers containing SPS sites respond to a given level of NICD more effectively. They do not, however, appear to affect the amount of NICD required for their initial activation, i.e., the threshold required for the enhancer to be switched on. This implies that burst size modulation and response threshold can be uncoupled and potentially could be encoded independently in the DNA sequence.

Regional Factors Prime Enhancers for Fast and Sustained Activation

Under ectopic NICD conditions, m5/m8 and sim both produce sustained transcription profiles in the MSE and NE, whereas elsewhere they generate stochastic and “bursty” transcription. This suggests that other factors are “priming” the enhancers to respond to NICD. Good candidates are the factors involved in DV patterning at this stage, the bHLH transcription factor Twist (Twi) and/or the Rel protein Dorsal (dl) whose endogenous gradients reach the region where m5/m8 and sim generate sustained profiles in response to eve2-NICD (Figure S6B) (Zinzen et al., 2006b). Furthermore, m5/m8 and sim both contain Twist and Dorsal binding motifs (Figure S6A) and previous studies indicated that Twist is important for sim activity although it was not thought to regulate to m5/m8 (Zinzen et al., 2006a).

To test if Twist and Dorsal are responsible for the different dynamics of NICD-induced transcription in the MSE versus DE (Figure 3C), we mutated the Twist and/or Dorsal binding motifs in m5/m8 (Figure S6A). Strikingly, mutating the three Twist or two Dorsal motifs produced a delay in the start of transcription in both wild-type and eve2-NICD embryos. These effects were even more pronounced when Twist and Dorsal motifs were mutated together (Figure 6B). Thus, without Twist or Dorsal, m5/m8 requires a higher threshold of NICD for activation or responds more slowly to the same threshold. The mean transcription levels were also reduced in all cases (Figure 6A).

Mutating the Twist motifs had two additional effects: the overall proportion of active cells in the MSE was reduced (Figure 6C) and few of those exhibited the sustained profile observed with wild-type m5/m8 (Figures 6E and 6F). Instead, most displayed a “bursty” transcription profile (Figure 6E), similar to those elicited by NICD in the DE. Although the mutated Twist motifs led to bursty profiles in wild-type embryos, these effects were partially rescued when ectopic NICD was provided (Figures 6C, 6F, and S6C). When both Dorsal and Twist motifs were mutated, the proportions of active nuclei and of nuclei with sustained profiles decreased even in the presence of ectopic NICD (Figures 6C, 6F, and S6C). The decrease in the overall proportion of active cells suggests that Twist and Dorsal regulate the ability of m5/m8 to activate transcription in response to Notch. In agreement, in embryos with two alleles of a reporter, the proportion of cells transcribing from both alleles was much lower for m5/m8simtwi than for m5/m8 (Figure 6D). Additionally, in those nuclei where both reporters were active, there was considerably more variability in the onset times for m5/m8simtwi compared to m5/m8 (Figure S6E). The results are therefore consistent with a role for Twist and Dorsal in priming the m5/m8 enhancer to become active in response to Notch and produce sustained activity. In their absence, the ability of m5/m8 to initiate transcription becomes much more stochastic and resembles that of m8NE (Figures 6F and S6D).

DISCUSSION

Developmental signaling pathways have widespread roles, but currently we know relatively little about how signaling information is decoded to generate the right transcriptional outcomes. We set out to investigate principles that govern how Notch activity is read by target enhancers in the living animal, using the MS2-MCP system to visualize nascent transcripts in Drosophila embryos and focusing on two enhancers that respond to Notch activity in the MSE. Three striking characteristics emerge. First, MSE enhancers are sensitive to changes in the levels of NICD, which modulate the transcriptional burst size rather than increasing burst frequency. Second, the activation of both MSE enhancers is highly synchronous. Indeed, within one nucleus the two enhancers become activated within few minutes of one another. Third, both MSE enhancers confer a sustained response in the wild-type context. This synchronized and persistent activity of the MSE enhancers contrasts with the stochastic and bursty profiles that are characteristics of most other enhancers that have been analyzed (Little et al., 2013; Fukaya et al., 2016; Fritzsch et al., 2019) and relies on the MSE enhancers being “primed” by regional transcription factors Twist and Dorsal. We propose that such priming mechanisms are likely to be of general importance for rendering enhancers sensitive to signals so that a rapid and robust transcriptional response is generated.

Priming of Enhancers Sensitizes the Response to NICD

Transcription of most genes occurs in bursts interspersed with refractory periods of varying lengths that are thought to reflect the kinetic interactions of the enhancer and promoter (Bartman et al., 2016). However, the MSE enhancers appear to sustain transcription for 40–60 min, without detectable periods of inactivity, though very short off periods might not have been resolved by our assays. Calculation of the ACF in traces from these nuclei suggest very slow transcriptional dynamics (Figures S4E and S4D) (Desponds et al., 2016; Lammers et al., 2018), consistent with one long period of activity rather than overlapping short bursts. This fits with a model where promoters can exist in a permissive active state, during which many “convoy”s of polymerase can be fired without reverting to a fully inactive condition.
The rapid successions of initiation events are thought to require Mediator complex (Tantale et al., 2016), which was also found to play a role in the NICD-mediated increase in residence time of CSL complexes (Gomez-Lamarca et al., 2018). We propose that sustained transcription from m5/m8 and sim reflects a switch into a promoter permissive state, in which general transcription factors like Mediator remain associated with the promoter so long as sufficient NICD is present, allowing repeated re-initiation.

However, the ability to drive fast and sustained activation is not a property of NICD itself. For example, when ectopic NICD was supplied, cells in many regions of the embryo responded asynchronously and underwent short bursts of activity. Furthermore, variable and less sustained cell-by-cell profiles were generated in the MSE region when the binding motifs for Twist and Dorsal in m5/m8 were mutated. The presence of these regional factors appears to sensitize the enhancers to NICD, a process we refer to as enhancer priming. This has two consequences. First, it enables nuclei to rapidly initiate transcription in a highly coordinated manner once NICD reaches a threshold level. Second, it creates an effective “state transition” so that the presence of NICD can produce sustained activity (Figure 7). We propose a priming mechanism, rather than classic cooperativity, because Twist and Dorsal alone are insufficient to drive enhancer activity.

See also Figure S5.
Figure 6. Twist and Dorsal Prime the Response of m5/m8 to NICD

(A) Mean activity levels in wild-type (top) and eve2-NICD (bottom) when Twist and/or Dorsal binding motifs in m5/m8 are mutated.

(B) Heatmaps of mutated enhancer activity in MSE nuclei in the wild-type and eve2-NICD, sorted by onset time. Dashed lines indicate onset times for unmutated enhancer.

(C) Proportion of active cells in the MSE in wild-type and eve2-NICD embryos when Twist and/or Dorsal motifs are mutated, compared to unmutated m5/m8.

(D) Proportion of active cells transcribing two alleles at any point, in embryos containing two copies of m5/m8 or of m5/m8DtwiDdl.

(E) Examples of transcription traces from wild-type and mutated m5/m8 in MSE nuclei from the wild-type and eve2-NICD embryos. Profiles from m5/m8DtwiMSE cells exhibit “bursty” transcription. ON periods are marked with a gray line.

(F) Proportion of MSE cells per embryo displaying a sustained profile of transcription, defined by ≥ one burst of >10 min. Median, quartiles, and SD are shown. Gray lines and heatmaps are re-plotted from Figures 3G and 3H. n = 4 (m5/m8WT), 5 (m5/m8DtwiWT), 4 (m5/m8DdlWT), 4 (m5/m8Dtwi eve2-NICD), 3 (m5/m8Dtwi eve2-NICD), 3 (m5/m8Ddl eve2-NICD), 3 (m8NE WT), 5 (m8NE eve2-NICD), and 3 (m5/m8DtwiDdl x2) embryos.

See also Figure S6.
required prior to NICD recruitment, although both may continue to play a role after transcription is initiated, as suggested by the lower mean levels when Twist or Dorsal motifs are mutated. Another contributory factor may be recruitment of the co-repressor complex containing CSL and Hairless, whose presence at primed enhancers could poise for activation and set the threshold (Barolo and Posakony, 2002).

Our explanation that the synchronous activation of the MSE enhancers reflects their requirements for a critical concentration of NICD is borne out by their responses when levels of NICD are increased. Notably, while sim and m5/m8 had almost identical dynamics in wild-type embryos, their response to ectopic NICD differed, suggesting that they detect different thresholds. Indeed, doubling the dose of ectopic NICD further accelerated the onset time of sim in agreement with the model that the enhancers detect NICD levels. Threshold detection does not appear to rely on the arrangement of CSL motifs, as onset times of m5/m8 or sim were unaffected by changes in the spacing of CSL paired sites. In contrast, mutating Twist- or Dorsal-binding motifs in m5/m8 delayed the onset, arguing that these factors normally sensitize the enhancer to NICD, enabling responses at lower thresholds.

We propose that enhancer priming will be widely deployed in contexts where a rapid and consistent transcriptional response to signaling is important, as in the MSE where a stripe of cells with a specific identity is established in a short time window. In other processes where responses to Notch are more stochastic, as during lateral inhibition, individual enhancers could be preset to confer different transcription dynamics. This suggests that paired motifs can “use” the NICD present more efficiently, potentially because optimally configured sites increase the likelihood that at least one NICD will be bound at any time. Interestingly, even though m5/m8 and sim contain different arrangements and numbers of CSL motifs they have converged to produce the same mean levels of transcription in wild-type embryos.

NICD Regulates Transcription Burst Size

Manipulating NICD levels revealed that Notch-responsive enhancers act as analog devices that can measure and broadcast variations in levels. Increased NICD levels have a consistent effect on enhancer activity irrespective of the priming state of the enhancer: an increase in burst size. Transcriptional bursting has been formalized as a two-state model where the promoter toggles between ON and OFF states, conferring a transcription initiation rate when ON (Peccoud and Ycart, 1995; Larson et al., 2009). Changes in duration or frequency of bursts lead to an overall increase in transcription. Most commonly, differences in enhancer activity have been attributed to changes in switching-on probability (Kon), leading to changes in burst frequency (Larson et al., 2013; Senecal et al., 2014; Fukaya et al., 2016; Fritzsch et al., 2018; Lammers et al., 2018; Berrocal et al., 2018). We were therefore surprised to find that higher doses of NICD did not increase burst frequency. Instead, they produced bigger bursts, both by increasing bursting amplitude, equivalent to the rate of transcription initiation, and bursting length, indicative of the total time the enhancer stays in the ON state. Modifications to the CSL motifs also impacted on the same parameters. Thus, enhancers with paired motifs (SPS) produced larger transcription bursts than those where the motifs are further apart. This suggests that paired motifs can “use” the NICD present more efficiently, potentially because optimally configured sites increase the likelihood that at least one NICD will be bound at any time. Interestingly, even though m5/m8 and sim contain different arrangements and numbers of CSL motifs they have converged to produce the same mean levels of transcription in wild-type embryos.
Two models would be compatible with the observations that effective NICD levels alter the burst size. In the first model, increasing the concentration of NICD when the enhancer is activated would create larger Pol II clusters. This is based on the observation that low-complexity activation domains in transcription factors can form local regions of high concentration of transcription factors, so-called “hubs,” which in turn are able to recruit Pol II (Mir et al., 2017; Tsai et al., 2017; Lu et al., 2018). As the lifetime of Pol II clusters appears to correlate with transcriptional output (Cho et al., 2016), the formation of larger Pol II clusters would in turn drive larger bursts. In the second model, NICD would be required to keep the enhancer in the ON state, for example, by nucleating recruitment of Mediator and/or stabilizing a loop between enhancer and promoter, which would in turn recruit Pol II in a more stochastic manner. General factors such as Mediator have been shown to coalesce into phase-separated condensates that compartmentalize the transcription apparatus (Cho et al., 2018; Sabari et al., 2018; Boija et al., 2018), and these could form in a NICD-dependent manner. Whichever the mechanism, persistence of the clusters and/or ON state requires NICD yet must be compatible with NICD having a short-lived interaction with its target enhancers (Gomez-Lamarca et al., 2018). Furthermore, the fact that the activity of m5/m8 saturates with one eve2-NICD construct and cannot be enhanced by providing a more active promoter suggests that there is a limit to the size or valency of the clusters that can form.

Although unexpected, the ability to increase burst size appears to be a conserved property of NICD. Live imaging of transcription in response to the Notch homolog, GLP-1, in the C. elegans gonad also shows a change in burst size depending on the signaling levels (Lee et al., 2019, this issue). As the capability to modulate burst size is likely to rely on the additional factors recruited, the similarities between the effects in fly and worm argue that a common set of core players will be deployed by NICD to bring about the concentration-dependent bursting properties.

STAR METHODS

Enhancer regulation of transcriptional bursting parameters revealed by forced chromatin looping. Mol. Cell 62, 237–247.

Berrocal, A., Lammers, N.C., Garcia, H.G., and Eisen, M.B. (2018). Kinetic sculpting of the seven stripes of the Drosophila even-skipped gene. bioRxiv. https://doi.org/10.1101/335901.

Bischof, J., Björklund, M., Furger, E., Schertel, C., Taipale, J., and Basler, K. (2013). A versatile platform for creating a comprehensive UAS-ORFeome library in Drosophila. Development 140, 2434–2442.

Bischof, J., Maeda, R.K., Hediger, M., Karch, F., and Basler, K. (2007). An optimized transgenesis system for Drosophila using germ-line-specific phiC31 integrases. Proc. Natl. Acad. Sci. USA 104, 3312–3317.

Blackwood, E.M., and Kadonaga, J.T. (1998). Going the distance: a current view of enhancer action. Science 281, 60–63.

Blythe, S.A., and Wieschaus, E.F. (2016). Establishment and maintenance of heritable chromatin structure during early Drosophila embryogenesis. Elife 5, 1–21.
Boja, A., Klein, I.A., Sabari, B.R., Dall’Agnese, A., Coffey, E.L., Zamudio, A.V., Li, C.H., Shrinivas, K., Manteiga, J.C., Hanner, N.M., Abraham, B.J., et al. (2018). Transcription factors activate genes through the phase-separation capacity of their activation domains. Cells, 1–14.

Boije, H., Rulands, S., Duddczig, S., Simons, B.D., and Harris, W.A. (2015). The independent probabilistic firing of transcription factors: A paradigm for clonal variability in the zebrafish retina. Dev. Cell 34, 532–543.

Bothma, J.P., Garcia, H.G., Esposito, E., Schlissel, G., Gregor, T., and Levine, M. (2014). Dynamic regulation of eve stripe 2 expression reveals transcriptional bursts in living Drosophila embryos. Proc. Natl. Acad. Sci. USA 111, 10598–10603.

Bothma, J.P., Norstad, M.R., Alamos, S., and Garcia, H.G. (2018). LlamaTags: a versatile tool to image transcription factor dynamics in live embryos. Cell 173, 1810–1822.e16.

Bray, S.J. (2006). Notch signalling: a simple pathway becomes complex. Nat. Rev. Mol. Cell Biol. 7, 678–689.

Chang, H.H., Hemberg, M., Barahona, M., Ingber, D.E., and Huang, S. (2008). Transcriptome-wide noise controls lineage choice in mammalian progenitor cells. Nature 453, 544–547.

Cho, W.K., Jayanth, N., English, B.P., Inoue, T., Andrews, J.O., Conway, W., et al. (2018). Activation of the Notch signaling pathway in vivo elicits changes in CSL nuclear dynamics. Dev. Cell 44, 611–623.e7.

Cho, W.K., Spiller, D.G., Mullins, J.J., Rand, D.A., Davis, J.R., et al. (2011). Dynamic analysis of stochastic transcription cycles. PLoS Biol. 9, e1000607.

Kopczynski, C.C., Atton, A.K., Fechtel, K., Kooch, P.J., and Muskatchev, M.A. (1988). Delta, a Drosophila neurogenic gene, is transcriptionally complex and encodes a protein related to blood coagulation factors and epidermal growth factor of vertebrates. Genes Dev. 2, 1723–1735.

Kosman, D., and Small, S. (1997). Concentration-dependent patterning by an ectopic expression domain of the Drosophila gap gene Knirps. Development 124, 1343–1354.

Kramatschek, B., and Campos-Ortega, J.A. (1994). Neuroectodermal transcription of the Drosophila neurogenic genes Espl and HLH-m5 is regulated by proneural genes. Development 120, 815–826.

Lammers, N.C., Galistyan, V., Reimer, A., Medin, S.A., Wiggins, C.H., and Garcia, H.G. (2018). Binary transcriptional control of pattern formation in development. bioRxiv. https://doi.org/10.1101/335919.

Larson, D.R., Fritzsche, C., Sun, L., Meng, X., Lawrence, D.S., and Singer, R.H. (2013). Direct observation of frequency modulated transcription in single cells using light activation. Elife 2, 1–20.

Larson, D.R., Singer, R.H., and Zenklusen, D. (2009). A single molecule view of gene expression. Trends Cell Biol. 19, 630–637.

Lee, C., Shin, H., and Kimble, J. (2019). Dynamics of Notch-dependent transcriptional bursting in its native context. Dev. Cell 50, this issue, 426–435.

Little, S.C., Tikhonov, M., and Gregor, T. (2013). Precise developmental gene expression arises from globally stochastic transcriptional activity. Cell 154, 789–800.

Lu, H., Yu, D., Hansen, A.S., Ganguly, S., Liu, R., Heckert, A., Darzacq, X., and Zhou, Q. (2018). Phase-separation mechanism for C-terminal hyperphosphorylation of RNA polymerase II. Nature 558, 318–323.

Mir, M., Reimer, A., Haines, J.E., Li, Y.Y., Stadler, M., Garcia, H., Eisen, M.B., and Darzacq, X. (2017). Dense bicoid hubs accentuate binding along the morphogen gradient. Genes Dev. 31, 1784–1794.

Mishchenko, Y. (2015). A fast algorithm for computation of discrete Euclidean distance transform in three or more dimensions on vector processing architectures. SIVP 9, 19–27.

Morel, V., Le Borgne, R., and Schweisguth, F. (2003). Snail is required for Delta endocytosis and Notch-dependent activation of single-minded expression. Dev. Genes Evol. 213, 65–72.

Morel, V., and Schweisguth, F. (2000). Repression by Suppressor of Hairless and activation by Notch are required to define a single row of single-minded expressing cells in the Drosophila embryo. Genes Dev. 14, 377–388.

Nam, Y., Sliz, P., Pear, W.S., Aster, J.C., and Blacklow, S.C. (2007). Cooperative assembly of higher-order Notch complexes functions as a switch to induce transcription. Proc. Natl. Acad. Sci. USA 104, 2103–2108.

Nandagopal, N., Santat, L.A., LeBon, L., Sprinzak, D., Bronner, M.E., and Elowitz, M.B. (2018). Dynamic ligand discrimination in the Notch signaling pathway. Cell 172, 869–880.e19.

Ng, K.K., Yui, M.A., Mehta, A., Siu, S., Irwin, B., Pease, S., Hirose, S., Elowitz, M.B., Rothenberg, E.V., and Kueh, H.Y. (2018). A stochastic epigenetic switch controls the dynamics of T-cell lineage commitment. Elife 7, 1–38.

Padovan-Merhar, O., Nair, G.P., Biash, A.G., Mayer, A., Scarfone, S., Foley, S.W., Wu, A.R., Churchman, L.S., Singh, A., and Raj, A. (2015). Single mammalian cells compensate for differences in cellular volume and DNA copy number through independent global transcriptional mechanisms. Mol. Cell 58, 339–352.

Peccoud, J., and Ycart, B. (1995). Markovian modeling of gene-product synthesis. Theor. Popul. Biol. 48, 222–234.

Raj, A., and van Oudenaarden, A. (2008). Nature, nurture, or chance: stochastic gene expression and its consequences. Cell 135, 216–226.

Raser, J.M., and Shea, E.K.O. (2006). Control of Stochasticity in eukaryotic gene expression Jonathan. Science 304, 1811–1814.

Sabari, B.R., Dall’Agne, A., Boja, A., Klein, I.A., Coffey, E.L., Shrinivas, K., Abraham, B.J., Har nett, N.M., Zamudio, A.V., Manteiga, J.C., et al. (2018).
Coactivator condensation at super-enhancers links phase separation and gene control. Science 361, eaar3958.

Senecal, A., Munskey, B., Proux, F., Ly, N., Braye, F.E., Zimmer, C., Mueller, F., and Darzacq, X. (2014). Transcription factors modulate c-Fos transcriptional bursts. Cell Rep. 8, 75–83.

So, L.H., Ghosh, A., Zong, C., Sepúlveda, L.A., Segev, R., and Golding, I. (2011). General properties of transcriptional time series in Escherichia coli. Nat. Genet. 43, 554–560.

Tantale, K., Mueller, F., Kozulic-Pirher, A., Lesne, A., Victor, J.M., Robert, M.C., Capozi, S., Chouaib, R., Bäcker, V., Mateos-Langerak, J., et al. (2016). A single-molecule view of transcription reveals convoys of RNA polymerases and multi-scale bursting. Nat. Commun. 7, 12248.

Tsai, A., Muthusamy, A.K., Alves, M.R., Lavis, L.D., Singer, R.H., Stern, D.L., and Crocker, J. (2017). Nuclear microenvironment modulates transcription from low-affinity enhancers. Elife 6, 158–160.

Vassin, H., and Campos-Ortega, J.A. (1987). Genetic analysis of Delta, a neurogenic gene of Drosophila melanogaster. Genetics 116, 433–445.

Wernet, M.F., Mazzoni, E.O., Çelik, A., Duncan, D.M., Duncan, I., and Desplan, C. (2006). Stochastic spineless expression creates the retinal mosaic for colour vision. Nature 440, 174–180.

Xu, H., Sepúlveda, L.A., Figard, L., Sokac, A.M., and Golding, I. (2015). Combining protein and mRNA quantification to decipher transcriptional regulation. Nat. Methods 12, 739–742.

Zabidi, M.A., Arnold, C.D., Schernhuber, K., Pagani, M., Rath, M., Frank, O., and Stark, A. (2015). Enhancer-core-promoter specificity separates developmental and housekeeping gene regulation. Nature 518, 556–559.

Zinzen, R.P., Cande, J., Ronshaugen, M., Papatsenko, D., and Levine, M. (2006a). Evolution of the ventral midline in insect embryos. Dev. Cell 11, 895–902.

Zinzen, R.P., Senger, K., Levine, M., and Papatsenko, D. (2006b). Computational models for neurogenic gene expression in the Drosophila embryo. Curr. Biol. 16, 1358–1365.
**STAR METHODS**

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Experimental Models: Organisms/Strains** | | |
| *D. melanogaster* His2Av-mRFP; nos-MCP-eGFP | Bloomington Drosophila Stock Center | RRID:BDSC_60340 |
| *D. melanogaster* nos-MCP-eGFP (II) | Bloomington Drosophila Stock Center | RRID:BDSC_63821 |
| *D. melanogaster* His2Av-RFP (III) | Bloomington Drosophila Stock Center | RRID:BDSC_23650 |
| *D. melanogaster* Df(3R)DlFX3 | Vassil and Campos-Ortega, 1987 | N/A |
| *D. melanogaster* Neur[11] | Bloomington Drosophila Stock Center | RRID:BDSC_2747 |
| *D. melanogaster* TTG (TM3, twi-Gal4, UAS-2xeGFP, Sb, Ser) | Bloomington Drosophila Stock Center | RRID:BDSC_6663 |
| *D. melanogaster* Ovo-FLP | Bloomington Drosophila Stock Center | RRID:BDSC_8727 |
| *D. melanogaster* betaTub85D-FLP | Bloomington Drosophila Stock Center | RRID:BDSC_7196 |
| *D. melanogaster* Vas-int; attP40 | Bischof et al., 2007 | N/A |
| *D. melanogaster* vas-int; attP-51D | Bloomington Drosophila Stock Center | RRID:BDSC_24483 (RFP cre-ed out) |
| *D. melanogaster* m5/m8-peve-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m5/m8-hsp70-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m5/m8-pm5-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m5/m8-pm6-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m5/m8-pm7-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m5/m8-pm8-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m5/m8-psimE-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* sim-peve-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* sim-psimE-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m8NE-peve-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* simSPS-peve-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m5/m8insSPS-peve-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m5/m8ATAT-peve-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m5/m8ADat-peve-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m5/m8ATADat-peve-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* m5/m8-peve-MS2-lacZ (II) | This study | N/A |
| *D. melanogaster* eve2-NICD (II, 25C) | This study | N/A |
| *D. melanogaster* eve2-NICD (II, 51D) | This study | N/A |
| **Oligonucleotides** | | |
| Oligonucleotides for cloning of enhancers and promoters in MS2 reporters and mutagenesis | This study | Table S1 |
| **Recombinant DNA** | | |
| pLacZ2-attB | Bischof et al., 2013 | N/A |
| pCR4-24XMS2SL-stable | Addgene | RRID:Addgene_31865 |
| 22FPE (eve.st.2-Pme) | Kosman, and Small, 1997 | N/A |
| pGEM-t-easy | Promega | N/A |
| pattB | Bischof et al., 2013 | N/A |
| pMT-NICD | Bray lab | N/A |
| **Software and Algorithms** | | |
| Tracking code | This study | https://github.com/juliafs93/FryEmbryo3DTracking |
| Modelling changes in transcription | This study | https://github.com/juliafs93/FFR_ACF |
| Matlab 2018a | Mathworks | N/A |
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Sarah J. Bray (sjb32@cam.ac.uk).

Experimental Animals

*Drosophila melanogaster* flies were grown and maintained on food consisting of the following ingredients:

Glucose 76g/l, Cornmeal flour 69g/l, Yeast 15g/l, Agar 2.5ml/l. Embryos were collected on apple juice agar plates with yeast paste. Animals of both sexes were used for this study.

Cloning and Transgenesis

**Generation of MS2 Reporter Constructs**

MS2 loops were inserted upstream of a lacZ transcript within the 5'UTR and then the resulting reporter was combined with different enhancers and promoters. 24 MS2 loops were cloned from pCR4-24XMS2SL-stable (Addgene #31865) into pLacZ2-attB (Bischof et al., 2013) using Eco RI sites. The m5/m8, sim and m8NE enhancers (Zinzen et al., 2006a; Kramatschek and Campos-Ortega, 1994) were amplified from genomic DNA and cloned into pattB-MS2-LacZ using HindIII/Agel sites (primers in Table S1). Subsequently the promoters hsp70, peve, pm5, pm6, pm7, pm8 and psimE were cloned by Gibson Assembly (Gibson, 2011) in pattB-m5/m8-MS2-LacZ, pattB-sim-MS2-LacZ and/or pattB-m8NE-MS2-LacZ (primers in Table S1) using the AgeI restriction site and incorporating a Eagl site.

Su(H), Twi, dI and Sna binding motifs were identified using ClusterDraw2 using the PWM from the Jaspar database for each transcription factor. Motifs with scores higher than 6 and p values < 0.001 were selected. Primers to create incorporating a m5/m8-MS2-LacZ affinity bicoid To generate Expression of Ectopic NICD and recombined with repository (https://benchling.com/juliafs/f_/AyerQ4a4-dynamics-notch-transcription-paper/).

**Expression of Ectopic NICD**

To test expression from two MS2 reporters in each cell (Figures 2 and S2) His2Av-RFP (BDSC #23650) was recombined with nos-MCP-GFP (BDSC #60340) in the third chromosome and combined with m5/m8-peve, sim-peve or m5/m8Δwi Δaf-m8-peve MS2 reporters. m5/m8-peve x2, sim-peve x2 and m5/m8Δwi Δaf-m8-peve x2 embryos were obtained from the stocks m5/m8-peve-MS2-LacZ ; His2Av-RFP, nos-MCP-GFP, sim-peve-MS2-LacZ ; His2Av-RFP, nos-MCP-GFP, respectively; while m5/m8-peve + sim-peve embryos were obtained from crossing sim-peve-MS2-LacZ ; His2Av-RFP, nos-MCP-GFP females with m5/m8-peve-MS2-LacZ males.
To observe transcription from MS2 reporters in conditions of ectopic Notch activity the FRT-STOP-FRT cassette had to be first removed from the eve2-NICD construct by expression of a flippase in the germline. To do so flies containing ovo-FLP (BDSC #7196) females were crossed with others containing eve2-FRT-STOP-FRT-NICD, His2Av-RFP and nos-MCP-GFP. The female offspring of this cross (ovo-FLP/+ ; eve2- FRT-STOP-FRT-NICD/+ ; His2Av-RFP, nos-MCP-GFP) induced FRT removal in the germline and were crossed with the MS2 reporters to obtain embryos expressing ectopic NICD. We note that only half of the embryos present the eve2-NICD chromosome, which could be distinguished by ectopic MS2 activity and an ectopic cell division of all the cells in the eve2 stripe after gastrulation. The other 50% embryos obtained from this cross were used as the wild type controls. This strategy was used to observe transcription from m5/m8-peve, sim-peve, m8NE-peve, m5/m8-pm5, simSPS-peve, m5/m8tw-peve, m5/m8at-peve and m5/m8at-peve. To measure transcription from 2eve2-NICD (Figures 4 and S4) removal of the FRT-STOP-FRT cassette was induced from the male germline to avoid recombination. To do so, betaTub85D-FLP (BDSC #7196) females were crossed with 2xeve2-NICD males and the male offspring of this cross (betaTub85D-FLP/Y ; 2xeve2-NICD/+), which induces FRT removal in the germline, were crossed with m5/m8-peve-MS2-lacZ ; His2AvRFP, nos-MCP-GFP or simpleve-MS2-lacZ ; His2AvRFP, nos-MCP-GFP females. As in the previous strategy, only half of the embryos presented the 2eve2-NICD chromosome and were distinguished by the ectopic activity. To express two m5/m8-peve reporters in conditions of ectopic NICD activity, m5/m8-peve and eve2-NICD were recombined in the second chromosome and embryos were obtained by crossing m5/m8-peve-MS2-lacZ ; His2AvRFP, nos-MCP-GFP females with betaTub85D-FLP/Y ; m5/m8-peve, eve2-NICD/+ males. Embryos were selected by the presence of two MS2 spots in each cell, which also ectopically expressed NICD.

METHOD DETAILS

Live Imaging

Embryos were dechorionated in bleach and mounted in Voltalef medium (Samaro) between a semi-permeable membrane and a coverslip. The ventral side of the embryo was facing the coverslip in all videos except when looking at transcription in the DE region, for which they were mounted laterally. Videos were acquired in a Leica SP8 confocal using a 40x apochromatic 1.3 objective and the coverslip. The ventral side of the embryo was facing the coverslip in all videos except when looking at transcription in the DE region.

QUANTIFICATION AND STATISTICAL ANALYSIS

Image Analysis

Videos were analyzed using custom Matlab (Matlab R2018a, Mathworks) scripts (available at https://github.com/juliafs93/FryEmbryo3DTracking)

Briefly, the His2Av-RFP signal was used to segment and track the nuclei in 3D. Each 3D stack was first filtered using a median filter, increasing the contrast based on the profile of each frame to account for bleaching and a fourier transform log filter (Garcia et al., 2013). Segmentation was performed by applying a fixed intensity threshold, 3D watershed accounting for anisotropic voxel sizes (Mishchenko, 2015) to split merged nuclei and thickening each segmented object. Nuclei were then tracked by finding the nearest object in the previous 2 frames which was closer than 6 μm. If no object was found, that nuclei was kept with a new label, and only one new object was allowed to be tracked to an existing one. After tracking, the 3D shape of each nucleus in each frame was used to measure the maximum fluorescence value in the GFP channel, which was used as a proxy of the spot fluorescence. We note than when a spot cannot be detected by eye this method detects only background, but the signal:background ratio is high enough that the subsequent analysis allows to classify confidently when the maximum value is really representing a spot.

In experiments with two MS2 reporters the maximum intensity pixel per nucleus does not allow to separate transcription from the two alleles. To do so, the 3D Gaussian spot detection method from (Garcia et al., 2013) was implemented in the existing tracking, such that each spot was segmented independently and associated with the overlapping nuclei. In this manner only active transcription periods were detected and no further processing of the traces was required.

MS2 Data Processing

From the previous step we obtained the fluorescent trace of each nuclei over time. Only nuclei tracked for more than 10 frames were retained. First nuclei were classified as inactive or active. To do so the average of all nuclei (active and inactive) was calculated over time and fitted to a straight line. A median filter of 3 was applied to each nuclei over time to smooth the trace and ON periods were considered when fluorescent values were 1.2 times the baseline at each time point. This produced an initial segregation of active (nuclei ON for at least 5 frames) and inactive nuclei. These parameters were determined empirically on the basis that the filters retained nuclei with spots close to background levels and excluded false positives from bright background pixels. The mean
fluorescence from MCP-GFP in the inactive nuclei was then used to define the background baseline and active nuclei were segregated again in the same manner. The final fluorescence values in the active nuclei were calculated by removing the fitted baseline from the maximum intensity value for each, and normalizing for the percentage that the MCP-GFP fluorescence in inactive nuclei decreased over time to account for the loss of fluorescence due to bleaching. Nuclei active in cycles before nc14 were discarded based on the timing of their activation.

In all videos, time into nc14 was considered from the end of the 13th syncytial division. When this was not captured the videos were synchronized by the gastrulation time. Plots showing mean fluorescent levels were obtained by calculating the mean and SEM of all fluorescent traces for multiple embryos aligned by the beginning of nc14. Calculating the mean levels of multiple embryos taken individually returned very similar profiles, indicating there is little embryo-to-embryo variability. In Figures 1C and 3C the total mRNA production per cell (in AU) was calculated by adding all the normalized fluorescent intensities for each nucleus.

Each embryo was classified into the 4 regions (ME, MSE, NE and DE) by drawing rectangular shapes in a single frame and finding which centroids overlapped with each region. In eve2-NICD these regions along the DV axis were defined within the eve2 stripe (~6-7 cells wide in all videos). In wild type embryos ME and MSE regions were drawn in the whole field of view (~150x150 μm anterior half of the embryo).

**Definition of Bursting Properties**

Bursts were defined as periods the median-filtered signal was higher than 1.2 times the baseline for at least 5 frames within a period from 15 min into nc14. These defined the burst duration and the time off between bursts. The amplitude was defined as the mean value within each burst period. The proportion of active cells was defined as the percentage of cells that switch on at any point after 15 min in each of the defined regions. 'Sustained' transcription was defined as nuclei with at least one burst longer than 10 min. This was based on analyzing regions where separated burst of activity were detected (mesoderm and dorsal ectoderm) where most bursts were <10 min. Off periods shorter than circa 2 mins would not have been resolved because the MS2 loops were positioned within the 5'UTR and the limit of resolution depends on the time taken for a Pol II molecule to complete transcription.

Onsets and ends of transcription were defined as the beginning of the first burst and the end of the last respectively (also starting at 15 min into nc14). In Figures 2 and S2 to be more precise in measuring the onsets and end points of transcription for both MS2 alleles they were scored manually as the first and last frame a spot is detected and randomly assigned 'allele 1' or 'allele 2'. The total variability was the variance of all onsets or end points, combining both alleles. The extrinsic variability was calculated as the covariance of onsets and ends between alleles 1 and 2. The remaining (total - covariance) corresponds to the intrinsic variability within each cell.

**Statistical Analysis**

In figure legends, n number indicates number of embryos imaged for each biological condition. Where appropriate, n number next to heatmaps indicates total number of cells combining all embryos for each biological condition. Plots showing mean levels of transcription and SEM (standard error of the mean) combine all traces from multiple embryos from the same biological condition. Violin plots show the bursting properties (amplitude, burst duration and off period) for each independent burst in all traces in multiple embryos, therefore the n number can be significantly greater than the number of cells in each condition. Because these properties do not follow a normal distribution, their statistical significance was tested with two Kolmogorov-Smirnov test. Levels of significance are indicated in the figure legends.

**Modelling Changes in Kinetic Parameters of Transcription**

We used a two-state promoter model of transcriptional activation in which the promoter switches between OFF and ON with constants \( K_{on} \) and \( K_{off} \) and releases mRNAs at a rate \( r \) when the promoter is ON (Figure 4E). This model also accounts for the residence time of polymerase on DNA while transcribing the gene (the elongation time \( T \)), so it is capturing what the MS2 system detects, ie. the number of nascent mRNA on the gene, rather than overall levels of mRNA in the cell. We take as a starting point expressions from (Choubey et al., 2015) for the mean and variance of the number of nascent mRNAs \( \langle m \rangle \) in steady state:

\[
\langle m \rangle = \frac{r T K_{off}}{K_{on} + K_{off}} \quad (1)
\]

\[
\text{Var}(m) = \langle m \rangle \left[ 1 + \frac{2 r K_{off}}{(K_{on} + K_{off})^2} + \frac{2 r K_{off}}{(K_{on} + K_{off})^3} \left( e^{-T} K_{off} - 1 \right) \right] \quad (2)
\]

We take the elongation time, \( T \), to be fixed for a given gene. Thus, according to Equation 1, the levels of transcription could increase in three ways: by increasing \( r \), increasing \( K_{on} \) or decreasing \( K_{off} \).

Thus, because of this degeneracy, observing a change in \( \langle m \rangle \) is alone insufficient to determine which underlying bursting parameter is being tuned to drive that change. However, we can make progress by incorporating the intrinsic noise of transcription into our analysis, since Equation 2 indicates that changes to bursting parameters that have equivalent effects on the mean may
nonetheless lead to different noise signatures. To do this, we calculate the Fano factor, which is defined as the variance divided by the mean:

\[Fano(m) = \frac{\text{Var}(m)}{\langle m \rangle}\] (3)

\[= \frac{1 + \frac{2rK_{\text{off}}}{(K_{\text{on}} + K_{\text{off}})^2}}{1 + \frac{2K_{\text{off}}}{(K_{\text{on}} + K_{\text{off}})^2} \left( e^{\frac{T(K_{\text{on}} + K_{\text{off}})}{T}} - 1 \right)}\] (4)

Where we see that the expression for the Fano factor is identical to the quantity inside the brackets in Equation 2.

Next, we examine how changes to each bursting parameter in turn will affect the Fano factor and Mean, respectively, demonstrating how these signatures can be used to uncover the drivers of observed changes between different experimental conditions.

**Pol II Initiation Rate \((r)\)**

We start by considering the case when \(r\) is modulated. In the discussions that follow, we assume a situation in which we are comparing two experimental conditions that exhibit observable differences in their mean rate of expression, \(\langle m \rangle\):

\[\alpha(m_1) = \langle m_2 \rangle\] (5)

Our goal is to determine whether the modulation of specific parameters corresponds reliably with changes in the mean and Fano factor. To do this, we undertake analysis of the functional form of the partial derivatives of these empirical measures with respect to each parameter.

From Equation 1, we have:

\[\frac{\partial \langle m \rangle}{\partial r} = \frac{TK_{\text{on}}}{\kappa}\] (6)

\[\frac{\partial \langle m \rangle}{\partial r} > 0\] (7)

Where, for convenience, we have introduced the shorthand \(\kappa = K_{\text{on}} + K_{\text{off}}\). So we see that \(\langle m \rangle\) is monotonic with \(r\): an increase in \(r\) always leads to an increase in the mean (and vice versa). The strict inequality applies because the right-hand-side of Equation 6 can be zeros if no expression occurs. For the Fano factor, we have:

\[\frac{\partial \text{Fano}}{\partial r} = \frac{2K_{\text{off}}}{\kappa^2} \left( 1 + \frac{e^{-\tau T - 1}}{\kappa T} \right)\] (8)

\[\frac{\partial \text{Fano}}{\partial r} \geq 0\] (9)

Unlike the mean, it is possible that a change in \(r\) could lead to no observable modulation in the Fano factor; however, this only holds for exceptionally small values of \(\kappa T\). More importantly, we see that it is impossible for the Fano factor to decrease when \(r\) is increased.

Thus, we conclude that an increase in \(r\) must coincide with an increase in both the mean rate of expression and in the Fano factor, i.e. the ratio between the Fano factors \(Fano(m_2)\) and \(Fano(m_1)\) where \(\langle m_2 \rangle = \alpha(m_1)\) would always be greater than 1 (Figure S3D, top panel).

**Activation Rate \((Kon)\)**

As with \(r\), we begin by examining how \(\langle m \rangle\) changes in response to a change in \(Kon\):\n
\[\frac{\partial \langle m \rangle}{\partial Kon} = \frac{rT}{\kappa} - \frac{rTK_{\text{off}}}{\kappa^2}\] (10)

\[= \frac{rT}{\kappa} \left( 1 - \frac{K_{\text{on}}}{\kappa} \right)\] (11)

\[\frac{\partial \langle m \rangle}{\partial Kon} \geq 0\] (12)

Thus, as with \(r\), the mean rate of expression increases monotonically in response to increases in \(Kon\). Next, for the Fano factor, we have:

\[\frac{\partial \text{Fano}}{\partial Kon} = 2rK_{\text{off}} \left( -\kappa^3 (2 + e^{-\tau}) + \frac{3\kappa^4}{T} (1 - e^{-\tau}) \right)\] (13)
To gain further insight, we need to examine limiting cases for the quantity $\kappa T$, which encodes the relative magnitude of the elongation time and switching rates, and which dictates the noise characteristics of the system.

We start with the case where $\kappa T << 1$:

$$\frac{\partial \text{Fano}}{\partial K_{\text{off}}} = -\frac{2rK_{\text{off}}}{\kappa^3} \left( 2 + 1 - \kappa T - \frac{3(1 - \kappa T - 1)}{\kappa T} \right)$$  \hspace{1cm} (15)

$$= -\frac{2rK_{\text{off}}}{\kappa^3} (3 - \kappa T - 3)$$  \hspace{1cm} (16)

$$= -\frac{2rK_{\text{off}}}{\kappa^3} (0)$$  \hspace{1cm} (17)

$$= -\frac{2rK_{\text{off}}}{\kappa^3} (3 - \kappa T - 3)$$  \hspace{1cm} (18)

$$\frac{\partial \text{Fano}}{\partial K_{\text{on}}} = 0$$  \hspace{1cm} (19)

For the opposite limit, where $\kappa T >> 1$, we have:

$$\frac{\partial \text{Fano}}{\partial K_{\text{on}}} = -\frac{2rK_{\text{off}}}{\kappa^3} \left( 2 + 0 - \frac{3(1 - 0)}{\kappa T} \right)$$  \hspace{1cm} (20)

$$= -\frac{4rK_{\text{off}}}{\kappa^3}$$  \hspace{1cm} (21)

$$\frac{\partial \text{Fano}}{\partial K_{\text{off}}} \leq 0$$  \hspace{1cm} (22)

So we see that, an increase in $\langle m \rangle$ that is driven by an increase in $K_{\text{on}}$ will coincide with a decrease in the Fano factor. Thus, unlike $r$, where the signs of the change in the mean and Fano factor are the same, we find that the signs of the changes in the mean and Fano factor are opposite in the case of changes driven by $K_{\text{on}}$, i.e., the ratio between the Fano factors $\text{Fano}(m2)$ and $\text{Fano}(m1)$ where $\langle m2 \rangle = \alpha \langle m1 \rangle$ would always be smaller than 1 (Figure S3D, middle panel).

**Off Rate ($K_{\text{off}}$)**

For the mean, we have:

$$\frac{\partial \langle m \rangle}{\partial K_{\text{off}}} = -\frac{rK_{\text{off}}}{\kappa^2}$$  \hspace{1cm} (23)

$$\frac{\partial \langle m \rangle}{\partial K_{\text{off}}} \leq 0$$  \hspace{1cm} (24)

Thus, as expected, an increase in $K_{\text{off}}$ leads to a decrease in $\langle m \rangle$. In keeping with our treatment in the case of $K_{\text{on}}$, we next examine the functional form of the Fano factor in the small and large $\kappa T$ limits. For $\kappa T << 1$, we expand about $\kappa T = 0$ to obtain an expression for the Fano factor:

$$\text{Fano} = 1 + \frac{2rK_{\text{off}}}{(K_{\text{off}} + K_{\text{off}})^2} + \frac{2rK_{\text{off}}}{(K_{\text{off}} + K_{\text{off}})^3} \left( \frac{1 - \kappa T - 1}{\kappa T} \right)$$  \hspace{1cm} (25)

$$= 1$$  \hspace{1cm} (26)

Thus, consistent with our findings for $K_{\text{on}}$, the Fano factor is largely insensitive to changes in $K_{\text{off}}$ for small $\kappa T$. This holds for $r$ as well, though we did not state so explicitly above. Next, we approximate the large $\kappa T$ limit by setting $e^{-\kappa T} = 0$:

$$\text{Fano} = 1 + \frac{2rK_{\text{off}}}{\kappa^2} + \frac{2rK_{\text{off}}}{\kappa^3} \left( \frac{0 - 1}{\kappa T} \right)$$  \hspace{1cm} (27)
removing the background baseline and normalizing for bleaching as described in the MS2 data processing section. When the onset of transcription was different between experiments (e.g., WT vs eve2-NICD) they were shifted to compare equivalent times. The Fano factor was calculated as the intrinsic variability divided by the mean over time:

\[ Fano = \frac{\sigma^2}{\langle m \rangle} \]  

Differentiating, we obtain:

\[ \frac{\partial Fano}{\partial K_{off}} = 2r \left( \frac{1}{\kappa^2} - \frac{2K_{off}}{\kappa} \right) \]  

\[ = \frac{2r}{\kappa^2} \left( 1 - \frac{2K_{off}}{\kappa} \right) \]  

The expression above reveals that, unlike \( r \) and \( K_{on} \), the direction of the change of the Fano factor in response to a change in \( K_{off} \) not fixed, but depends upon the relative sizes of \( K_{on} \) and \( K_{off} \), i.e., the ratio between the Fano factors \( Fano(m_2) \) and \( Fano(m_1) \) where \( \langle m_2 \rangle = \alpha \langle m_1 \rangle \) could be smaller or greater than 1 (Figure S3D, bottom panel). Numerical simulations confirm this result.

**Stochastic simulations**

We next tested with simulations whether the Fano factor ratio can be used as a diagnostic tool of the underlying changes in the mean. We used stochastic simulations of transcription based on the Gillespie algorithm (Gillespie 1976) of the same two-state promotor model but using additional parameters to more resemble the biological MS2 data (accounting for the time MS2 loops are detected, acquisition time and adding experimental noise, Figure S3F).

We then tested whether we could recover the same trends in Fano factor ratios in the simulation as expected from the mathematical model (Figure S3D). However, given that changes in \( K_{off} \) can produce Fano factor ratios greater or smaller than 1, calculation of the Fano factor and comparing whether it is greater or smaller than 1 alone is not sufficient to infer which parameter is being modified to produce the observed changes in the mean.

**Utilizing the Autocorrelation Function (ACF)**

The results of our analysis thus far indicate that modulations in \( r \) and \( K_{on} \) lead to distinct, well defined signatures in mean and Fano factors of experimentally observed expression levels. However, the degeneracy of the Fano factor shift with respect to changes in \( K_{off} \) necessitates the incorporation of an additional observable, if we are to be able to distinguish the underlying drivers of changes between experimental conditions. To this end, we utilize the empirical Autocorrelation Function of our experimental MS2 traces.

The ACF function provides information about the speed of the system and the elongation rate (Desponds et al., 2016; Lammers et al., 2018). Intuitively, the more rapid the time scale with which the system switches between activity states (the larger \( \kappa \) is), the faster the ACF decays. We used the same simulations to test if the autocorrelation function changes in different ways depending on the modified parameters, to help distinguishing between the 3 scenarios to increase the mean. If the dynamics are fast (Figure S3E, right column, \( K_{off} = 0.1 \) s\(^{-1} \) and \( K_{on} = 0.2 \) s\(^{-1} \)) no changes in the ACF were observed in any of the three cases. When the dynamics are slower (Figure S3E, left column, \( K_{on} = 0.01 \) s\(^{-1} \) and \( K_{off} = 0.02 \) s\(^{-1} \)), then the AC function shifts to the right (from \( \langle m_1 \rangle \) to \( \langle m_2 \rangle \)) when \( K_{off} \) decreases. No changes are observed when \( r \) or \( K_{on} \) increase.

Therefore looking at both the Fano factor ratio and the autocorrelation function (when the dynamics are slow enough), provides enough information to distinguish between the three ways in which the mean can change (Figure S3B):

- increase in \( r \): FFRatio > 1 and no change in ACF
- increase in \( K_{on} \): FFRatio < 1 and no change in ACF
- decrease in \( K_{off} \): FFRatio < 1 or > 1 and shift to the right in ACF

**Estimating Fano factor from empirical data**

When applied to real MS2 traces, raw fluorescence profiles from each cell were processed by applying a median filter of 3, removing the background baseline and normalizing for bleaching as described in the MS2 data processing section. When the onset of transcription was different between experiments (e.g., WT vs eve2-NICD) they were shifted to compare equivalent times. The Fano factor was calculated as the intrinsic variability divided by the mean over time:

\[ Fano = \frac{\sigma^2}{\langle m \rangle} \]  

\[ = \frac{Var(m) - CoVar(m)}{\langle m \rangle} \]  

The intrinsic component was calculated by subtracting an estimation of the extrinsic variability form the total noise. The contribution from the extrinsic noise, normally calculated from the covariance of two transcription traces from the same cell, was calculated
by using neighbouring nuclei as proxy of two loci in the same cell and calculating their covariance. Using the experiments where two MS2 reporters are present in each cell we validated the contribution from extrinsic noise is equivalent within cell and across neighbouring cells. Both FFRatio and ACF were calculated by doing 50 bootstraps of all available traces and calculating the mean and SD.

DATA AND CODE AVAILABILITY

Scripts for tracking and analysis of MS2 videos are available at https://github.com/juliafs93/FryEmbryo3DTracking. The code developed for the modelling approach to infer changes in parameters of transcription causing changes in mean levels of transcription is available at https://github.com/juliafs93/FFR_ACF.