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Novel Fluorescent and Photoconvertible Fusions Reveal Dorsal Activator Dynamics

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- Abstract Over the last two decades, new *in vivo* and *in cellulo* imaging technologies have
- ¹⁶ uncovered the inherently dynamic nature of transcriptional regulation in embryonic
- ¹⁷ development and, in particular, in the fruit fly *D. melanogaster*. These technologies have made it
- ¹⁸ possible to characterize the subnuclear and single-molecule dynamics of transcription factors.
- ¹⁹ However, a lack of appropriate fluorescent protein fusions has, until now, limited these studies to
- ²⁰ only a few of the dozens of important transcription factors in the fruit fly gene regulatory network
- ²¹ dictating early development. Here, we report the creation of four new fluorescent protein fusions
- to Dorsal, a member of the NF- κ B/Rel family that initiates dorsal-ventral patterning. We
- 23 generated and characterized two bright fluorescent protein fusions for Dorsal, meGFP and
- ²⁴ mNeonGreen, and two photoconvertible fluorescent protein fusions, mEos4a and mNeonGreen.
- ²⁵ We show that removal of the DsRed2 cassette commonly used to mark the CRISPR integration
- restores endogenous Dorsal mRNA and protein levels and enables the fusion allele to rescue a
- *dorsal* null allele, meeting the gold standard for endogenous function of a tagged protein in a
- ²⁸ fruit fly. We then demonstrate that our bright fluorescent protein fusions can be used to dissect
- ²⁹ the spatiotemporal dynamics of stable Dorsal clusters that traverse the nucleoplasm and
- ³⁰ uncovered that these clusters preferentially interact with active sites of Dorsal-modulated
- transcription. We further demonstrate that our photoconvertible fluorescent protein fusions
- make it possible to detect individual molecules of Dorsal in the nuclei of developing embryos.
- ³³ These new fluorescent protein fusions constitute a valuable resource for the community to
- elucidate the role of Dorsal activator dynamics in dictating fruit fly early embryonic development.
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1 Introduction

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- ³⁷ Over the last two decades, new *in vivo* and *in cellulo* microscopy technologies have made it possible
- 38 to uncover the dynamics of regulatory transcription factors as they interact with transcription sites
- ³⁹ to activate or repress gene expression in the context of embryonic development (*Wagh et al., 2023*;
- 40 Boka et al., 2021; Lu and Lionnet, 2021). The emerging picture is one where activators or repressors
- an only transiently occupy their target binding sites at enhancers (*Mir et al., 2017, 2018; Lu and Lion*
 - net, 2021; Donovan et al., 2019)—and sometimes act in the context of spatially localized hubs or

43 clusters (*Mir et al., 2017, 2018; Sabari et al., 2018; Wei et al., 2020; Kawasaki and Fukaya, 2023*)—to

⁴⁴ regulate the stochastic transcription process underpinned by transcriptional bursting (*Rodriguez*

and Larson, 2020; Lammers et al., 2020b; Leyes Porello et al., 2023; Meeussen and Lenstra, 2024).
 These discoveries have been partially fueled by an ever-increasing palette of fluorescent pro-

- teins, which are fused to transcription factors to enable direct measurements of their real-time
- 49 dynamics. This palette now includes fluorescent proteins that are suitable for a wide range of
- ⁴⁹ live imaging experiments: brighter and more photostable fluorescent proteins (meGFP, mClover3,
- 50 mNeonGreen, mStayGold) enable longer-term imaging, and photoactivatable (PA-GFP) and photo-
- s1 convertible (mEos3.2, Dendra2) fluorescent proteins enable superresolution and single-molecule
- ⁵² imaging (all reviewed in *Rodriguez et al.* (2017)).

Yet, despite this ever-growing toolbox of fluorescent proteins, it is time-consuming and challenging to fuse newly engineered fluorescent proteins to a protein of interest in a manner that preserves that protein's endogenous functionality (*Chen et al., 2011; Cranfill et al., 2016*). For example, it has proven particularly difficult to generate fluorescent protein fusions for early transcription factors (TFs) in the developing fruit fly (*Drosophila melanogaster*) embryo, such as Bicoid

⁵⁸ and Dorsal (*Reeves et al., 2012*; *Singh et al., 2022*).

⁵⁹ Dorsal, a transcriptional activator belonging to the NF-*κ*B/Rel family (*Hong et al., 2008; Gilmore,* ⁶⁰ **2006**), initiates fruit fly embryonic dorsal-ventral patterning via a maternally deposited concentra-⁶¹ tion gradient (*Hong et al., 2008; Gilmore, 2006*).Despite its crucial role in the developmental cas-⁶² cade of the early fruit fly embryo, studies of Dorsal dynamics have been limited by the availability ⁶³ of functional fluorescent protein fusions. Only a single fluorescent protein fusion. Dorsal-mVenus.

⁶⁴ meets the gold standard for maintaining endogenous Dorsal activator activity: a single copy of a

⁶⁵ Dorsal-mVenus transgene allele complements (or, "rescues" the development of) a Dorsal null al-

⁶⁶ lele (e.g. *dl[1]*). Subsequently, *Alamos et al.* (2023) successfully generated a Dorsal-mVenus CRISPR

⁶⁷ knock-in allele—using the same combination of linker and fluorescent protein as in the transgene

by **Reeves et al.** (2012)—that also rescues the *dl*[1] null allele.

While Dorsal-mVenus has proven exceedingly useful in measuring the nuclear levels and dorsoventral gradient of Dorsal in the embryo (*Reeves et al., 2012*; *Alamos et al., 2023*), we sought to

⁷¹ answer outstanding questions about the sub-nuclear dynamics of Dorsal, including measuring the

⁷² activity of subnuclear clusters and the binding dynamics of single Dorsal molecules. These specific

r3 experimental goals required a fluorescent protein more photostable than mVenus, as well as a

⁷⁴ photoconvertible fluorescent protein.

Here, we describe an expansion of the fluorescent protein palette for endogenous Dorsal, fea-75 turing four new fluorescent protein CRISPR knock-in fusions: the brighter and more photostable 76 meGFP (Cormack et al., 1996; Zacharias et al., 2002) and mNeonGreen (McKinney et al., 2009), and 77 the photoconvertible mFos4a (Kopek et al., 2017: Paez-Segala et al., 2015) and Dendra2 (Gurskava 78 et al., 2006). All four fusions produce viable progeny from females homozygous for the Dorsal-FP 79 allele, and we show that removing the DsRed marker cassette enables our Dorsal-mNeonGreen al-80 lele to rescue a Dorsal null allele. Thus, these new fusions are suitable for studying the endogenous 81 dvnamics of Dorsal. 82

⁸³ We demonstrate the potential of these brighter and more photostable meGFP and mNeon-

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- ⁸⁴ Green fusions to study recently discovered clusters of Dorsal concentration that have been sug-
- ⁸⁵ gested to play a role in the regulation of Dorsal target genes (Yamada et al., 2019). Our fusions
- make it possible to track the dynamics of these clusters, revealing that Dorsal clusters tend to be
- ar in closer proximity to target genes than non-target genes—a phenomenon that we explore more
- deeply in a pair of accompanying papers (*Fallacaro et al., 2025; Dima et al., 2025*). Additionally, we
- ⁸⁹ demonstrate how the photoconvertible mEos4a and Dendra2 fusions enabled us to track single
- ⁹⁰ molecules of Dorsal binding to DNA for the first time, finding that Dorsal spends only a few seconds
- ⁹¹ bound to the DNA, an observation consistent with the binding times of several other transcription
- ₉₂ factor in the fruit fly and beyond (Lammers et al., 2020b; Lu and Lionnet, 2021). These two ex-
- ⁹³ periments, only made possible by the new Dorsal fusion alleles, help increase our understanding
- 94 of the dynamic process of transcriptional activation and demonstrate that the four Dorsal fusions
- ⁹⁵ presented in this paper will constitute a valuable resource for the community.

96 2 Results

97 2.1 Generation of novel Dorsal fusions

We fused two fluorescent proteins, meGFP and mNeonGreen (McKinney et al., 2009), and four 98 photoconvertible proteins, Dendra2 (*Gurskava et al., 2006*), mEos3.2 (*Zhang et al., 2012*), mEos4a, qq and mEos4b (Paez-Segala et al., 2015; Kopek et al., 2017), in-frame to the C-terminus of the Dor-100 sal protein via a 6xGlycine (6G) linker using an existing CRISPR/Cas9 protocol (*Gratz et al.* (2015): 101 Alamos et al. (2023); Methods Section 4.1 and Table S1). The knock-in was marked by a 3xP3-102 DsRed2-SV40polA cassette, which drives the expression of the fluorescent protein DsRed in the 103 adult eves and ocelli, allowing for rapid screening of successfully transformed adults. This DsRed 104 cassette, as we will refer to it from now on, was flanked by a pair of 3' and a 5' PiggyBac transposon 105 sites to allow for scarless removal of DsRed. The DsRed cassette was placed downstream of the flu-106 orescent protein's stop codon (Gratz et al., 2015) and upstream of the endogenous 3' untranslated 107 region (3'UTR). We characterized several successful (as indicated by DsRed+ adults), independent 108 integrations of each CRISPR knock-in *dorsal (dl)* allele. 109

The introduction of the fluorophore sequence can interfere with the regulation, production, or 110 function of the protein to which it is fused, resulting in impaired downstream functions. As an ini-111 tial assessment of the function of these fusions, we tested whether females homozygous for the 112 Dorsal fusion alleles were fertile and able to generate viable progeny. In the case of a maternally-113 deposited transcription factor like Dorsal, the ability for females homozygous for the fusion alleles 114 to produce viable progeny indicates that the developmental functions of Dorsal are intact. Females 115 homozygous for the dl-6G-mEos4b-DsRed and dl-6G-mEos3.2-DsRed alleles were not fertile, generat-116 ing no viable progeny at room temperature (approximately 22°C). This outcome indicates that the 117 maternally deposited copies of these Dorsal fusion proteins do not function well enough to drive 118 development (*Table 1*). As a result, we did not proceed with any further characterization of the 119 mEos4b nor mEos3.2 lines. 120

In contrast, females homozygous for the dl-6G-meGFP-DsRed, dl-6G-mNeonGreen-DsRed, dl-6G-121 mEos4a-DsRed, and dl-6G-Dendra2-DsRed alleles vielded viable puppe at room temperature. These 122 results indicate that these Dorsal fusion proteins maintain some level of normal function during 123 development. However, very few pupae were produced by females homozygous for these four al-124 leles. Only females homozygous for the dl-6G-Dendra2-DsRed allele produced sufficient progeny to 125 be maintained as a stable, homozygotic line; females homozygous for the other three alleles—*dl*-126 6G-meGFP-DsRed, dl-6G-mNeonGreen-DsRed, and dl-6G-mEos4q-DsRed—produced too few progeny 127 to be maintained as stable, homozygotic line. Only the *dl-6G-Dendrg2-DsRed* allele produced suffi-128 cient progeny to be maintained as a stable, homozygotic line. Additionally, when females homozy-120 gous for these four alleles were kept at elevated temperatures (25°C), they no longer yielded pupae. 130

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suggesting that these fusion alleles compromise the robustness of the function of Dorsal during 131 development in the face of temperature changes. 132

As a more quantitative measure of fly line viability, we measured and compared the embryo 133 hatch rate across these four homozygous alleles (*Figure 1A*). We counted the percentage of em-134 bryos, laid by females homozygous for each allele, that successfully hatched into larvae after 36 135 hours (*Methods Section 4.4*). We compared these hatch rates to the wild-type (untagged) dl allele 136 (vw; + ;+, hereafter vw) and the dl-6G-mVenus-DsRed allele (Alamos et al., 2023). Embryos from 137 wild-type *dl* females had the highest hatch rate, 85%, followed closely by embryos from *dl-6G*-138 mVenus-DsRed females, which hatched at a 75% rate. None of the embryos from females with 139 the new Dorsal fusion alleles exceeded the hatch rate of the dl-6G-mVenus-DsRed allele. dl-6G-140 Dendrg2-DsRed had the highest hatch rate of the new alleles (55% and 60%, for two independent 141 integrations), followed by dl-6G-mEos4g-DsRed (15% and 20%) and dl-6G-meGFP-DsRed (5% and 20%). 142 *dl-6G-mNeonGreen-DsRed* exhibited the lowest embrvo hatch rates (2% and 5%). 143

To further characterize these fusions, we performed live imaging to assess the resulting Dorsal 144 protein expression pattern. All four alleles drive a Dorsal gradient that gualitatively matches the 145 expected endogenous expression pattern of Dorsal protein: high nuclear Dorsal levels along the 146 ventral midline decreasing to nuclear exclusion of Dorsal towards the dorsal side of the embryo 147 Figure 18-E. Thus, our results demonstrated, at least gualitatively, that the maternal Dorsal protein 148 was being translated and imported to the nuclei as expected. 149

Despite the qualitative agreement between the endogenous Dorsal gradient and the gradient 150 resulting from our Dorsal fusions, the poor viability of our new fusion fly lines (Table 1) led us to 151 question whether these lines were faithful reporters of endogenous Dorsal function. Specifically, 152 we hypothesized that the poor viability of our CRISPR dl alleles could have three causes: off-target 153 CRISPR/Cas9-induced mutations, interference from the protein sequence linker, and/or interfer-154 ence from the presence of the DsRed cassette in the 3'UTR. 155

2.2 Removal of DsRed cassette restores embryo viability of Dorsal fusion fly lines 156

We investigated and corrected for the three potential causes of the poor embryo viability in our 157 homozygous fusion allele fly lines, as hypothesized in the previous section. First, we removed 158 off-target CRISPR/Cas9-induced mutations in other essential genes via out-crossing. Such muta-159 tions could lead to non-viability, without implicating maternal Dorsal function. Out-crossing for 160 six to eight generations did not improve embryo viability (Supplemental Information Section S1 : 161 Figure S1). 162

Second, we altered the linker sequence between Dorsal and the fluorescent protein fusion. 163 The specific sequence of certain linkers may interfere with the regulation, folding, or function of 164 maternal Dorsal more than the sequence of others. We were unable to identify an alternative 165 linker sequence that improved the embryo viability of these fly lines (Supplemental Information 166 Section S2 : Table 1). 167

Third, we assessed the effect of the DsRed marker cassette in the 3'UTR. The DsRed cassette 168 is located between the stop codon of the fluorescent protein and the start of the endogenous 3' 169 untranslated region (3'UTR) sequence. While this position does not alter the Dorsal protein coding 170 sequence, it does modify the Dorsal mRNA sequence, particularly the position and sequence of its 171 3'UTR. 3'UTRs play a significant role in mRNA stability, mRNA localization, regulation of translation. 172 and protein-protein interactions (Szostak and Gebauer, 2013: Buxbaum et al., 2015: Andreassi and 173 Riccio, 2009; Kuersten and Goodwin, 2003; Mayr, 2019). The presence of the DsRed cassette in the 174 3'UTR could interfere with any of these important molecular functions, leading to altered Dorsal 175 protein expression levels and impacting downstream target genes. 176 To assess the effect of the DsRed cassette on viability, we removed this sequence from the line 177 carrying the *dl-6G-mNeonGreen-DsRed* allele using scarless removal via the PiggyBac transposase

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Figure 1. Generation and characterization of novel Dorsal fusion proteins. (A) Embryo hatch rates for embryos laid by females homozygous for various Dorsal fusion alleles, quantified as fraction of embryos that had hatched 36 hours after being laid. Embryo hatch rates for an untagged *dl* allele (from *yw* flies) served as a control. Hatch rates for each allele can also be found in *Table 1*. (**B-E**) Images of our homozygous-viable Dorsal fusion lines taken on laser scanning confocal microscope during later nuclear cycle 14 prior to gastrulation: (**B**) Dorsal-mNeonGreen, (**C**) Dorsal-meGFP, (**D**) Dorsal-mEos4a, and (**E**) Dorsal-Dendra2.

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(Nyberg and Carthew 2022) We then quantified and compared the viability of the allele with and 179 without the DsRed marker cassette. In contrast to embryos from females with two copies the 180 dl-6G-mNeonGreen-DsRed allele, embryos from females with two copies of the dl-6G-mNeonGreen-181 $\Delta DsRed$ allele were viable and could produce a stable stock population. Given this improvement 182 in embryo viability, we then tested our dl-6G-mNeonGreen- Δ DsRed allele for its ability to maintain 183 embryo viability as a single copy. The ability of a transgenic dl allele to complement a mutant null dl 184 allele-here, dl[4]—is considered the gold standard for demonstrating normal protein function. We 185 found that embryos from dl-6G-mNeonGreen- Δ DsRed / dl[4] females were viable, but embyros from 186 dl-6G-mNeonGreen-DsRed / dll41 females were not. Thus, our results indicate that the interruption 187 in the 3'UTR reduces embryo viability. 188

2.3 Changes in the *dorsal* 3'UTR modulate mRNA levels and nuclear protein con centration

¹⁹¹ To identify the molecular cause of this reduced embryo viability, we measured and compared the ¹⁹² *dl* mRNA levels, Dorsal protein pattern, nuclear Dorsal protein concentrations, and Dorsal-target ¹⁹³ gene expression produced by the *dl-6G-mNeonGreen* allele with and without the DsRed cassette ¹⁹⁴ present, hereafter referred to as *DsRed* and $\Delta DsRed$, respectively.

First, we quantified the effect of the DsRed cassette on d/mRNA production by measuring mRNA 195 levels using quantitative real-time PCR (gPCR) on embryos collected from homozygous females of 106 the DsRed and the Δ DsRed lines. We determined the relative mRNA levels with respect to vw em-197 bryos carrying a wild-type d/ allele (Figure 2A). The ratio of dl-6G-mNeonGreen-DsRed to wild-type 198 dl mRNA levels was 0.5±0.1 (mean \pm standard error of the mean). indicating that the presence of 199 the DsRed cassette reduced mRNA levels by half. In contrast, the ratio between dl-6G-mNeonGreen-200 $\Delta DsRed$ and wild-type d/ was found to be 1.0+0.2, indicating that the scarless removal of the DsRed 201 cassette restored d/ mRNA levels to wild-type. As a result, we concluded that the significant reduc-202 tion in mRNA levels due to the presence of the DsRed cassette in the 3'UTR of the *dl* allele was a 203 likely cause of the reduced viability of the fly lines carrying the dl-6G-mNeonGreen-DsRed allele. 204

Second, to determine the downstream impact of these reduced mRNA levels on Dorsal protein 205 levels and localization. we measured the Dorsal-mNeonGreen protein gradient along the dorsal-206 ventral axis during development. We imaged the cross-section of embryos from both heterozygous 207 dl-6G-mNeonGreen-DsRed / His2Av-RFP (Figure 2B) and heterozygous dl-6G-mNeonGreen- Δ DsRed / 208 His2Av-RFP females (Figure 2C), where the His2Av-RFP fluorescence signal was used for nuclear 200 segmentation. The Dorsal-mNeonGreen protein gradient is gualitatively similar in the DsRed and 210 ΔDsRed embryos, with high nuclear Dorsal-mNeonGreen levels in the ventral nuclei, lower nuclear 211 Dorsal-mNeonGreen levels in dorsal nuclei, and negligible cytoplasmic Dorsal-mNeonGreen lev-212 els (*Figure 2B-C*). To quantify the Dorsal nuclear concentration gradient along the embryo, we fit 213 a Gaussian function to the nuclear Dorsal-mNeonGreen signal across the full dorsal-ventral axis 214 and determined its width and amplitude (Methods Section 4.8: Liberman et al. (2009): Reeves et al. 215 (2012)). Although the width of the Dorsal gradient was similar for *DsRed* and $\Delta DsRed$ embryos in nu-216 clear cycle 13, we observed a slight difference in nuclear cycle 14 (*Figure 2D*). When we measured 217 the amplitude of the Dorsal gradient, we found that the *DsRed* embryos had approximately half the 218 nuclear Dorsal-mNeonGreen fluorescence in their ventral-most nuclei as compared to the $\Delta DsRed$ 219 embryos (Figure 2E-F). Similarly, we found that the ventral nuclei in DsRed embryos contained a 220 little more than half (approximately 53% in nuclear cycle 14) the absolute Dorsal-mNeonGreen 221 protein concentration than the $\Delta DsRed$ embryos, as measured by Raster Image Correlation Spec-222 troscopy (RICS) (Supplementary Information Section S3 : Methods Section 4.9). 223

Finally, as expected, the reduced levels of nuclear Dorsal protein in the DsRed embryos led to altered mRNA expression patterns in downstream, Dorsal-regulated genes (*Figure S3*). We found

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that only 11% of the embryos had a wild-type expression pattern of the Dorsal-activated gene, *snail* in the ventral nuclei (*Figure S3B(i)*), with the remaining embryos exhibiting either a significantly

²²⁷ in the ventral nuclei (*Figure S3B(i)*), with the remaining embryos exhibiting either a significantly ²²⁸ reduced or entirely absent *sna* pattern (*Figure S3B(ii-iii)*). We posit that the aberrant expression

- ²²⁸ reduced or entirely absent *sna* pattern (*Figure S3B(II-III)*). We posit that the aberrant expression ²²⁹ of *snail* and other gene expression patterns led to the low percentage of hatched embryos of this
- ²²⁹ of *snall* and other gene expression patterns led to the low percentage of hatched embryos of th
- and other transgenic lines (*Table 1*).

We conclude that the presence of the DsRed marker cassette in the Dorsal 3'UTR reduces the viability of the fly line by halving *dl* mRNA levels, which in turn leads to downstream reduction of Dorsal protein levels and severely altered gene expression of Dorsal-activated genes. We can restore the viability of these new fusion fly lines by scarless removal of the DsRed cassette from







(A) *dl* mRNA expression ratio, as measured by qPCR, for embryos from females carrying two copies of the *dl-6G-mNeonGreen-DsRed* (DsRed; pink) and the *dl-6G-mNeonGreen-\DeltaDsRed* (Δ DsRed; blue) alleles, relative to a wild-type *dl* allele. *dl-6G-mNeonGreen-DsRed*: 0.5±0.1; *dl-6G-mNeonGreen-\DeltaDsRed*: 1.0±0.2 (mean ± standard error of the mean). (**B-C**) Representative cross-section images of the Dorsal-mNeonGreen fluorescent protein gradient and measurement of gradient width, as indicated by the perpendicular colored lines, in (**B**) *dl-6G-mNeonGreen-DsRed* / His2Av-RFP embryos and (**C**) *dl-6G-mNeonGreen-\DeltaDsRed* / His2Av-RFP embryos. Scale bar is 50 µm. (**D**) Width of the Dorsal-mNeonGreen gradient, as extracted from a Gaussian fit, in DsRed and Δ DsRed embryos during nuclear cycles 13 and 14 (dark regions) and the intervening mitosis (light region). (**E**) Quantification of the normalized Dorsal-mNeonGreen nuclear fluorescence in ventral (solid lines) and dorsal (dashed lines) nuclei for DsRed (pink) and Δ DsRed (blue) embryos during nuclear cycles 13 and 14. Line values are the mean of five DsRed embryos and three Δ DsRed embryos; shaded regions are SEM. (**F**) Ratio of Dorsal-mNeonGreen nuclear fluorescence in Δ DsRed to DsRed embryos in the ventral (solid black line) and dorsal (dashed, brown line) nuclei.

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236 2.4 Photostable fluorescent proteins uncover subnuclear Dorsal cluster dynamics 237 and their proximity to sites of transcription

Recently, regions of high Dorsal concentration—clusters—were found in the vicinity of target genes
 (*Yamada et al., 2019*). These clusters were found to be correlated with an increase in the mean rate
 of transcription of Dorsal target genes. However, because these measurements were performed
 using fixed tissue techniques, how Dorsal cluster dynamics dictate transcriptional control remains
 unknown.

Using the existing Dorsal-mVenus CRISPR fusion (*Alamos et al., 2023*) line and live imaging on a confocal microscope, we uncovered relatively stable, submicron-sized clusters of high Dorsal concentration that move about the nucleoplasm (*Figure 3*A). These initial results confirmed that these clusters were not just an artifact of the original fixed embryo staining with anti-Dorsal antibody labeling that was used to discover them (*Yamada et al., 2019*).

Yet, despite our ability to capture the clusters using the Dorsal-mVenus fusion, these clusters were only observable under a high spatial resolution and excitation laser intensity. Additionally, we noted that the fast movemnet of the clusters in the nucleus required a frame rate of ~20 seconds to accurately capture their dynamics. These two live imaging requirements led to rapid photobleaching, within 2-3 minutes, due to the poor photostability of mVenus (*Figure 3*B).

Our two newly developed, more photostable Dorsal fusions, Dorsal-meGFP and Dorsal-mNeonGreen, enabled us to perform longer-term imaging of these Dorsal clusters. Indeed, clusters were clearly visible using both fusions (*Figure 3*C,D). Further, while the measurable fluorescence intensity was reduced to half of the starting intensity within four minutes for the Dorsal-mVenus line, the fluorescence intensity of the Dorsal-meGFP line remained above 80% of the starting intensity past six

minutes (*Figure 3*B). This increased photostability enabled us to image and track Dorsal clusters
 for the full length of a nuclear cycle, as shown by the comparison between cluster movies of the
 Dorsal-mVenus and the Dorsal-meGFP fusions shown in *Figure 3*E and F.

With the ability to visualize Dorsal clusters in real time, we sought to uncover how they interact with target genes to regulate gene expression. To make this possible, we simultaneously imaged the Dorsal-mNeonGreen clusters and with the transcriptional activity of a *snail* reporter construct a target of Dorsal—labeled using the MS2 system (*Bertrand et al., 1998; Garcia et al., 2013; Lucas et al., 2013*) over the course of an entire nuclear cycle (*Figure 4*A). Here, nascent mRNA molecules are labeled using mCherry, such that sites of nascent transcript formation are visible as fluorescent puncta.

To investigate how the dynamics of these Dorsal clusters might dictate *snail* gene expression dynamics, we sought to characterize the positions of all clusters in a nucleus and compare them to the positions of actively transcribing *snail* loci as shown schematically in *Figure 4*B, and as exemplified using representative images in *Figure 4*C and D. As the definition of a cluster is challenging due to their varying intensity and size, along with a high background intensity from nuclear Dorsal fluorescence, we adopted a segmentation approach based on contour level sets. The two highest contour heights were defined as the "cluster" levels (*Figure 4*E).

Using these contour-based cluster segmentation results, we calculated the pairwise distances between an actively transcribing, MS2-labelled locus and all the clusters detected in the same nucleus, at the same z-plane. Speficially, we quantified the distribution of pairwise cluster-locus distances for the Dorsal-target gene *snail* (*Figure 4*F). As a negative control to which to compare *snail*, we generated the same distance distribution for the non-target gene *hunchback* (*Figure 4*F).

We found that there is a population of clusters that are within 300 nm of actively transcribing *snail* loci. This population was not present near non-target *hunchback* loci. These results suggest that the population of clusters in the vicinity of the *snail* reporter is a subset of clusters that are preferentially associated with loci targeted by Dorsal. In companion papers posted alongside this

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- С D Α Dorsal-mVenus 5 µm В Ε fluorescence intensity, normalized to t=0 0.5 6 min 2 min 0 min F Dorsal-meGFP Dorsal-mVenus 0 2 0 6 4 time from start of imaging (min) 0 min 2 min 6 min
- work, we leverage these reagents to further explore the dynamics of these clusters and their role
 in regulating transcriptional activity (*Fallacaro et al., 2025*; *Dima et al., 2025*).

Figure 3. Dorsal-meGFP and Dorsal-mNeonGreen fusions enable live imaging of Dorsal clusters. (A) Snapshots from movies of Dorsal-mVenus revealing clusters of high concentration (white arrows). **(B)** Dorsal-mVenus clusters showed rapid bleaching as compared to the minimal bleaching exhibited by Dorsal-meGFP when imaged on the same microscope, under the same imaging conditions. **(C, D)** Dorsal clusters are clearly visible (white arrows) in the context of fusions to (C) mNeonGreen and (D) meGFP. **(E, F)** Representative snapshots of the field of view used to quantify the total fluorescence intensity for each time point quantified in (B) for (E) Dorsal-mVenus and (F) Dorsal-meGFP.

286 285 Photoconvertible fluorescent proteins uncover Dorsal single-molecule dynam 287 ics

²⁸⁸ The recent development of lattice light sheet microscopy (*Chen et al., 2014*) has made it possible

- ²⁸⁹ to quantify the binding dynamics of transcription factors—such as the maternally-deposited tran-
- ²⁹⁰ scription factor Bicoid (*Mir et al., 2017*) and the uniformly distributed pioneer-like transcription
- ²⁹¹ factor Zelda (*Mir et al., 2018*)—in living, developing fruit fly embryos. However, single molecule
- ²⁹² detection of abundant transcription factors such as Bicoid, Zelda and Dorsal on a lattice light sheet

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Figure 4. Dorsal clusters preferentially associate with transcriptional loci. (**A**) Two sample nuclei illustrating that clusters are often spatially associated with active transcriptional loci. In the top montage, nucleus A, a *snail* transcription locus turns on seemingly within a Dorsal cluster and the two remain tightly associated for approximately two minutes until the Dorsal cluster dissolves into the nuclear Dorsal background. In the bottom montage, nucleus B, a Dorsal cluster emerges from the background pool of nuclear Dorsal and moves to associate with an already-active *snail* transcription locus. The two remain associated for approximately one minute until the locus turns off, while the Dorsal cluster remains visible. (**B**) Cartoon illustrating the measurement of pairwise distances between an actively-transcribing locus labeled with MS2 and the Dorsal clusters in a nucleus. (**C-D**) Example snapshots of the same nucleus with (C) an actively-transcribing *snail* labeled by MCP-mCherry and (D) a Dorsal-mNeonGreen cluster that is nearby. (**E**) Contour level sets used to bin the Dorsal-mNeonGreen intensity are shown for the example nucleus from (D), with the second highest contour level, deemed the "Dorsal cluster" contour, outlined in red. The position of the *snail*-MS2-mCherry spot from (C) is marked by a red dot. (**F**) Distribution of pairwise distances between all Dorsal clusters in a nucleus.

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microscope requires a fluorescent protein with two key properties: high signal-to-noise ratio (SNR) and sparse labeling. For these reasons, the existing Dorsal-mVenus fusion was not a suitable choice

²⁹⁵ for single-molecule measurements.

Our two new photoconvertible Dorsal fusions, Dorsal-mEos4a and Dorsal-Dendra2, solve both of these challenges, with improved SNR over Dorsal-mVenus and intrinsic photoconvertible properties, allowing for the measurement of the *in vivo* binding dynamics of individual Dorsal molecules. As a proof-of-concept, we imaged these new fusion under a custom MOSAIC microscope in the lattice light sheet imaging modality (*Chen et al., 2014*).

Excitation with a 488 nm laser enables bulk measurements of the unconverted Dorsal-mEos4a 301 (Figure 5A.C: Figure 54A.C) and Dorsal-Dendra2 (Figure 54E.G) fusion proteins in the ventral nuclei 302 of fruit fly embryos, where Dorsal is the most highly concentrated. We then photoconverted a 303 subpopulation of the Dorsal fusion proteins with a low power of a 560 nm laser and imaged the 304 resulting photoconverted Dorsal-mEos4a (Figure 5B: Figure 54B) and Dorsal-Dendra2 (Figure 5F) 305 proteins using 560 nm excitation. Specifically, we imaged using an exposure of 500 ms, which is 306 expected to blur out the contribution of freely diffusion Dorsal molecules and make it possible to 307 detect those molecules that are instead stably bound to the DNA (*Mir et al., 2017, 2018*). 308

The resulting photoconverted subpopulation of Dorsal-mEos4a was sparse enough to identify 309 single molecules of the Dorsal fusions bound to the DNA of individual nuclei (*Figure 5D: Figure S4D*). 310 These single molecules had high SNR and were photostable enough to track for up to eight sec-311 onds (*Figure 5D*: *Figure S4D*). The photoconverted subpopulation of Dorsal-Dendra2 was initially 312 too high, resulting in an SNR that was too low for accurate single-molecule tracking (*Figure 5*F). To 313 achieve single-molecule tracking over the same time-scale as Dorsal-mEos4a, we needed to first 314 image under 560 nm excitation for approximately four minutes (Figure S4F) to bleach most of the 315 photoconverted molecules. Only then was the unbleached, photoconverted subpopulation small 316 enough to achieve the required SNR and sparsity to track for a similar length of time (*Figure S4*H) 317 as the Dorsal-mEos4a molecules. Thus, our new photoconvertible fluorescent Dorsal fusions are 318 an ideal substrate to carry out single-molecule measurements of the binding dynamics of this tran-319 scription factor, as well as how these dynamics dictate output transcriptional dynamics of target 320 genes. 321

322 **3 Discussion**

Leveraging the advancing biochemical and optical properties of newly engineered fluorescent pro-323 teins can open new scientific avenues and help answer previously unresolved questions. However, 324 the rate-limiting step in adopting these fluorescent proteins—particularly for essential targets like 325 early developmental transcription factors—is achieving functional fusion with the protein of inter-326 est without disrupting its normal biological activity. In this study, we selected a papel of fluorescent 327 proteins with useful properties for longitudinal imaging and single molecule tracking experiments. 328 such as increased brightness, increased photostability, and the ability to photoswitch. We char-320 acterized their effects on function when fused to the maternal transcription factor Dorsal in the 330 fruit fly embryo, and demonstrated their potential to uncover biological phenomena that were 331 previously inaccessible. 332

We identified four fluorescent proteins (meGFP, mNeonGreen, mEos4a, Dendra2) that can be successfully fused to Dorsal while maintaining the ability of the Dorsal-fluorescent protein allele to function when present in two copies. However, these fusions did not fully rescue embryonic variability as homozygous alleles, let alone as heterozygotes combined with a *dorsal* mutant allele. We discovered that the DsRed marker commonly used to determine the successful CRISPRmediated knock-in of sequences into the genome, and not the nature of the fluroescent protein or the linker between this protein and Dorsal, was the culprit behind the loss of viability of our

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Dorsal fusions. Indeed, both mRNA and protein levels were significantly reduced by the presence 340 of the DsRed cassette in the 3'UTR of the dorsal gene. Upon removal of this cassette, both mRNA 341 and protein levels recovered to wild-type levels and embryonic viability was restored. The mecha-342 nism by which the presence of DsRed decreases dorsal mRNA levels remains to be uncovered. We 343 speculate that, due to the alteration of the 3'UTR, such reduction likely results from an alteration 344 of the *dorsal* mRNA lifetime. Regardless, these results serve as a reminder to the fly community 345 that, although convenient for the purposes of tracking alleles, markers such as the DsRed cassette 346 might significantly compromise the very developmental processes we seek to characterize. 347 Having characterized these fusions, we demonstrated their potential for shedding light on the 348

mechanisms of Dorsal action in two contexts. First, we showed how the increased photosability of our novel Dorsal-mNeonGreen and Dorsal-meGFP fusions made it possible to track subnuclear clusters of Dorsal protein within the nucleus. Second, we demonstrated the feasibility of using our fusions of Dorsal to photoactivatable fluorescent proteins mEos4a and Dendra2 for the characterization of Dorsal DNA binding at the single-molecule level.

For most developmental genes studied to date, the exact nature and timing of the molecular 354 mechanisms that underlie regulation of their transcription, and the nature of the bursts by which 355 this transcription is characterized still remain elusive (Rodriguez and Larson, 2020; Lammers et al., 356 2020b; Leyes Porello et al., 2023; Meeussen and Lenstra, 2024). The growing body of evidence 357 that sequence-specific and general transcription factors exist in transient clusters—also referred 358 to as hubs, microenvironments, or condensates—of high local concentration (Mir et al., 2017, 2018; 359 Tsai et al., 2017: Sabari et al., 2018: Cho et al., 2018, 2016: Zamudio et al., 2019: Klosin et al., 2020) 360 has provided a tantalizing new way of thinking about how these molecular mechanisms could play 361 out, as well as a potential avenue for measuring correlations with transcriptional bursting. How-362 ever, so far it has been challenging to establish the functional role of these clusters in regulating 363

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364 transcription (*McSwiggen et al., 2019*).

Using our Dorsal fusions to photostable fluorescent proteins, we imaged its clusters in space 365 and time and discovered that, on average, Dorsal clusters are more likely to be found in the vicinity 366 of its target gene *sngil* when compared to its non-target gene *hunchback*. In parallel collaborative 367 work, we have used the reagents presented here to further study the interaction between Dorsal 368 clusters and the transcriptional dynamics of Dorsal target genes (Fallacaro et al., 2025; Dima et al., 369 2025). Thus, our new Dorsal-mNeonGreen and Dorsal-meGFP fusions provide an ideal tool to study 370 the spatiotemporal underpinnings of Dorsal nuclear dynamics and the role these dynamics play in 371 dictating transcriptional control. 372

Finally, our new fusions of Dorsal to photoactivatable proteins made it possible to, for the first time, detect individual molecules of this important fly transcription factor as they bind and unbind from the DNA. Our proof of concept promises to make it possible to measure the Dorsal dwell time on the DNA as it has been previously done for Bicoid and Zelda (*Mir et al., 2017, 2018*), and to relate the localization and dynamics of this binding to the regulation of Dorsal target genes.

To sum up, our new fusions of Dorsal to novel fluorescent proteins will make it possible to reveal the mechanistic underpinnings of Dorsal and its regulation of transcription. We envision that these protein fusions will become a valuable resource for the fly community in performing the quantitative experiments necessary to reach a predictive understanding of cellular decision making in development.

383 4 Methods and Materials

384 4.1 Plasmids

To generate the Dorsal fluorescent protein knock-in alleles, we used a previously published CRISPR/Cas9 protocol (*Gratz et al., 2015; Alamos et al., 2023*). The CRISPR donor plasmids were modified from a *dl-6G-mVenus-DsRed* donor plasmid (*Alamos et al., 2023*). The donor plasmid carries the following insertion sequence: a 6xGlycine (6G) linker followed by the mVenus protein coding sequence, a stop codon, a 3xP3-DsRed2-SV40polA cassette flanked by PiggyBac transposon sites, and the endogenous 3'UTR sequence. The whole insertion sequence is flanked by two ~ 1 kb homology arms that target the endogenous Dorsal stop codon.

To generate the new fusion plasmids, the mVenus sequence was replaced with the desired flu-392 orescent protein sequence (meGFP, mNeonGreen, mEos3.2, mEos4a, mEos4b, or Dendra2), while 393 all other elements were left unmodified. Alternative linker plasmids were also generated for two of 394 the fluorphore fusions. mEos3.2 and mNeonGreen. The 6G linker (GGGGGG) of the *dl-6G-mEos3.2* 395 plasmid was replaced by three alternative linkers: 6G-10GS (GGGGGGGGGGGGGGGSGSGS), 6G-helix 396 (GGGGGGMSKGEEL) the MSKGEEL portion is the N-terminal helix of mVenus) and LL (LongLinker 397 SGDSGVYKTRAOASNSAVDGTAGPGSTGSS: a gift from Michael Stadler). The 6G linker of the dl-6G-398 mNeonGreen plasmid was replaced by the LL linker. All alternative linker sequences were generated 300 via gene synthesis by GenScript (Riiswijk, Netherlands), Cloning and sequencing to confirm the final 400 plasmid sequence was performed by GenScript, Inc. (Riiswiik, Netherlands). 401

For the synthetic guide RNA (sgRNA) plasmid, we used the previously published plasmid, *pU6-DorsalgRNA1*, which expresses a synthetic guide RNA (sgRNA or gRNA) (GUUGUGAAAAAGGUAU-UACG) that targets a sequence in the C-terminus of Dorsal on Chromosome 2 (*Alamos et al., 2023*). The list of all plasmids described in the current study can be found in *Table S2*, and full se-

quences for all plasmids can be accessed at https://benchling.com/garcialab/f_/THClp5A3-dorsalfusions-manuscript/.

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408 4.2 Transgenic Fly Lines

Each fluorescent protein fusion donor plasmid was co-injected with the pU6-DorsalgRNA1 plasmid 409 into embryos expressing Cas9 under the control of vasa regulatory sequences in the ovary (BDSC 410 #51324) by BestGene, Inc. (Chino Hills, CA, USA). Surviving adults were crossed to either w1118 411 or vw stocks and their offspring were screened for DsRed fluoresence in the adult eves. Transfor-412 mants were balanced with CvO by crossing to vw; Sp/CvO; + to generate a stable lines. Several 413 independent integrations were established as stable lines. The CRISPR insertions were confirmed 414 by PCR using primers recognizing the left homology arm. TS-F (GAGGGCGACAAAGGCAAAGA) and 415 Donor-R (CGCCACCACCTGTTCCTGT), and the right homology arm. Donor-F (GGGCAGCTTCACTC-416 CTTTCT) and TS-R (TACGCCGCACTAACGAATCT). The 3xP3-DsRed2-SV40polA eve marker cassette 417 was initially left in all lines to allow for simplified crossing schemes to other transgenic lines. 418

To generate the vw: Dorsal-meGEP-DsRed, eNosx2-MCP-mCherry / CvO : + and vw: Dorsal-meGEP-419 DsRed, eNosx2-MCP-mCherry / CvO; + transgenic lines were generated by recombining vw; Dorsal-420 meGFP-DsRed / CvO : + or vw: Dorsal-mNeonGreen-DsRed / CvO : + (Table S1) with vw: eNosx2-MCP-421 mCherry @VK22 / CvO; +, Female progeny lacking the CvO balancer (i.e. curly wings) were crossed 422 to vw: Sp/CvO: + males and the progeny of that cross were screened for individuals that had both 423 the DsRed and white+ markers visible in the adult eve, as well as the CvO balancer (curly wings) 424 present. Secondary, confocal microscopy screening of the embryos of the resulting stable lines 425 was done to confirm that the lines expressed both mCherry and meGFP or mNeonGreen in the 426 embryo. 427

The full list of fly lines described in this study can be found in *Table S1*.

429 4.3 Outcrossing

 $_{430}$ *yw* ; *Dorsal-Fluorescent Protein-DsRed / CyO* ; + females were crossed to *yw* males. The female $_{431}$ progeny were screened for DsRed expression and DsRed+ flies were then crossed again to *yw* $_{432}$ males. This outcrossing was done for 6-8 generations. The final generation of female progeny was $_{433}$ re-balanced with *CyO*, by crossing to *yw* ; *Sp/CyO* ; + to re-generate a stable line.

434 4.4 Embryo hatch test

To test for maternal Dorsal function in our Dorsal-FP CRISPR knock-in fusion lines, we measured embryo hatching rates for embryos laid by non-virgin females that carry two copies of the Dorsal-FP CRISPR fusion allele (i.e. homozygotes). Homozygotes were selected by screening for an absence of the balancer chromosome *CyO* and curly wings. Males were a mix of heterozygous and homozygous flies, which means that these embryo hatch tests were *not* an accurate indicator of zygotic Dorsal function, as the embryos themselves could either be heterozygous (one untagged, wildtype *dorsal* allele and one *dorsal*-FP allele) or homozygous (two *dorsal*-FP alleles). The following step-by-step protocol was used for each embryo hatch test:

- Prepare cages with females homozygous for Dorsal-FP-DsRed and mixed homozygous and
 heterozygous males from the same line.
- 2. Transfer embryos to a juice agar plate in rows each containing 5 or 10. Record the total
 number of embryos mounted, aiming for at least 100 embryos per hatch test.
- A447
 3. Place a small dab of yeast paste at one edge of the plate so the larvae crawl to it after they
 A448
 A447
- 449 4. Place small agar plate in a larger, covered, petri dish and leave for ~ 36 hours at room tem-450 perature (~ $22^{\circ}C$).
- 451
 5. Count the number of empty chorions (eggshells) to determine how many embryos are "hatched",
 452
 452
 and count the number of embryos still in their chorion to determine the number of "un-
- 453 hatched" embryos.

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6. Confirm that the number of hatched embryos matches the difference between the number
 of embyros mounted and the number of unhatched embryos.

Each round of hatch tests included an agar plate of embryos from a cage of *yw* females and

⁴⁵⁷ males and another agar plate of embryos from a cage of *yw*; *Dorsal-6G-mVenus-DsRed*;+ females

458 and males, both of which served as positive controls and points of comparison across different

⁴⁵⁹ biological replicates and days.

460 Embryo hatch tests were performed both prior to and after outcrossing.

461 4.5 DsRed removal

487

488

⁴⁶² DsRed marker was removed using PiggyBac transposase following an established protocol (*Nyberg* ⁴⁶³ *and Carthew, 2022*). The fly stock having PiggyBac transposase transgene used for the crosses is ⁴⁶⁴ w[1118]; Herm3xP3-ECFP, α tubuling-piggyBacK10M6 (BDSC 32070). The removal of DsRed was ⁴⁶⁵ confirmed by checking the absence of DsRed expression in the fly eyes as well as by PCR checks ⁴⁶⁶ using genomic DNA of each fly line as templates. To ensure the genomic edits for the FP of interest

467 is intact, PCR checks were performed and embryos from each fly line were imaged using primers

- DsRed: ATGGCCTCCTCCGAGGACGT and CTACAGGAACAGGTGGTGGC
- mNeonGreen: GTGAGCAAGGGCGAGGAGGA and CTTGTACAGCTCGTCCATGC
- mEGFP: GTGAGCAAGGGCGAGGAGCT and CTTGTACAGCTCGTCCATGC

• Dendra2: AACACCCCGGGAATTAACCT and CCACACCTGGCTGGGCAGGG

• mEos4a: GTTAGTGCGATTAAGCCAGA and TCGTCTGGCATTGTCAGGCA

473 4.6 Quantitative real-time PCR (qPCR)

To measure the mRNA levels driven by our different Dorsal fusions, embryos were collected from 474 homozygous females of both types (with DsRed, Δ DsRed) as well as wild-type fly line *yw* as control. 475 Grape juice plates were changed twice separated by one hour. The plates were then kept in the 476 cages for 45 minutes before embryo collection. The embryos were washed from the plates to a 477 mesh and dechorionated using 100% bleach for 3 minutes. The embryos were then washed with 478 deionized water to remove residual bleach. Only the embryos younger than no 13 were selected 479 and washed with 500 µl PBT in a microcentrifuge tube. RNA isolation was performed using standard 480 a TRIzol- based extraction method (Invitrogen, USA). The extracted RNA was treated with DNase 481 I (Thermo Scientific, USA) to remove genomic DNA and purified using the Monarch RNA Cleanup 482 Kit (New England Biolabs, USA) to remove any residual salts from the DNase buffer. 170 ng of the 483 collected RNA was reverse transcribed using the ProtoScript II First Strand cDNA Synthesis Kit (New 484 England Biolabs, USA) following the standard protocol, *actin* (act) was used as the reference gene. 485 The primer pairs used were: 486

- dl: TGG CTT TTC GCA TCG TTT CCA G and TGT GAT GTC CAG GGT ATG ATA GCG
- αctin: CCG TGA GAA GAT GAC CCA GAT C and TCC AGA ACG ATA CCG GTG GTA C

Genomic DNA was amplified using the same PCR primers and used as template for five 10-fold 489 serial dilutions to generatw the standard curve for the gPCR. The cDNA from with DsRed, Δ DsRed, 490 vw were used as templates for gPCR using the *dl* and *actin* primer sets described above. All the re-491 actions were performed in triplicates. The gPCR was performed using iTag Universal SYBR Green 492 Supermix (Bio-Rad, USA) in the CFX Opus 96 Real-Time PCR System (Bio-Rad, USA). The quantifica-493 tion cycle, C_a were determined using CFX Maestro Software (Bio-Rad, USA). The calibration curve 494 was generated by plotting the average C_a of the triplicate vs logarithm of initial template concen-495 tration for the serial dilutions. The PCR efficiency (E) of each primer pair was determined from the 49F slope of the calibration curve (Bustin et al., 2009) by calculating 407

 $E = 10^{\frac{-1}{slope}} - 1 \tag{1}$

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⁴⁹⁸ Finally, the relative gene expression ratio, *R*, is calculated by doing

$$R = \frac{(E_{dl})^{C_q^{yw} - C_q^{sample}}}{(E_{actin})^{C_q^{yw} - C_q^{sample}}}$$
(2)

where E_{dl} and E_{actin} are the PCR efficiencies for the *dl* and *actin* primer pairs, respectively, and C_q^{yw}

and C_{q}^{sample} are the quantification cycle for the control line *yw* and the sample (either with DsRed or

⁵⁰¹ ΔDsRed), respectively (*Pfaffl, 2001, 2004*):.

502 4.7 Live imaging

⁵⁰³ Embryos from females having one copy of *dl-mNeonGreen-DsRed* and one copy of *His2Av-RFP* as well

as embryos from females having one copy of *dl-mNeonGreen-\Delta DsRed* and one copy of *His2Av-RFP*

were collected. The *His2Av-RFP* transgene was obtained from the Bloomington Stock Center ([*]; Pw[+mC]=His2Av-mRFP1IL2. BDSC:23651). The embryos were dechorionated in 100% bleach for

⁵⁰⁶ Pw[+mC]=His2Av-mRFP1II.2. BDSC:23651). The embryos were dechorionated in 100% bleach for ⁵⁰⁷ 30 seconds and washed with deionized water to remove residual bleach. The embryos at nuclear

⁵⁰⁸ cycle 10 were selected manually.

509 4.8 Measurement of the Dorsal gradient

The embryos were end-on mounted in 1% low melt agarose, with the anterior pole touching the cov-510 erslip. The cross-section was imaged 150 µm from the anterior pole from nuclear cycle 10 until gas-511 trulation in 30-second time intervals with a frame time of 10 s. An LD C-Apochromat 40x/1.1 W Corr 512 objective was used. Analysis of the Dorsal gradient was done following previously published pro-513 tocols (Liberman et al., 2009; Reeves et al., 2012). Briefly, nuclear segmentation was performed 514 based on the His2Av-RFP signal. The amplitude of the gradient representing the Dorsal amount 515 in the ventral-most nuclei and the basal level representing the non-zero amount of Dorsal in the 516 dorsal-most nuclei were determined by Gaussian-fitting to the nuclear DI-mNeongreen signal. 517

518 4.9 RICS

519 Embryos were mounted on their ventral side manually. A C-Apochromat 40x/1.2 W autocorr objec-

tive was used. The images were collected at a 5x zoom (pixel size of 31.95 nm) and a frame time of

521 5.06 s. The analysis was performed following previously published protocols (*Schloop et al., 2024*;

522 Al Asafen et al., 2024; Dima and Reeves, 2024).

523 4.10 Embryo collection and preparation

Non-virgin, homozygous females and males from each Dorsal-FP line were crossed together in a cage. Virgin females from a line expressing a maternally-deposited TF and the MS2 coat proteins (e.g. *yw; Dorsal-mVenus-DsRed, pNos-MCP-mCherry; Dorsal-mVenus, pNos-MCP-mCherry, His2Av-iRFP*) were crossed to males carrying an MS2 reporter reporter gene.

Cages were fed a paste of dry, activated yeast mixed with water, which was placed on a petri dish lid containing grape juice agar and changed out at least once per day. Fly cages were allowed to lay for 90 to 120 minutes prior to embryo collection. Embryos were then mounted on microscopy slides in Halocarbon 27 Oil (Sigma-Aldrich, H8773) in between a #1.5 glass coverslip and a membrane semipermeable to oxygen (Lumox film, Starstedt, Germany) as described in *Garcia et al.* (2013) and *Bothma et al.* (2018).

534 4.11 Imaging

535 Data collection for Dorsal gradient assessment was performed on a Leica SP8 scanning confocal

microscope (Leica Microsystems, Biberach, Germany). Each Dorsal-FP fusion was excited at 488 nm

and its fluorescent signal detected by a Hybrid Detector (HyD) set to photon counting mode with a

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- spectral window of 496-546 nm. Average laser powers were not quantified, but were substantially
- ⁵³⁹ higher for mEos4a, mEos3.2, and Dendra2 as compared to meGFP and mNeonGreen to compen-
- sate for the formers' poor quantum yield in their green states. Pinhole was set to 1.0 Airy units
- ⁵⁴¹ (AU) for an emission peak wavelength of 509 nm. All data was taken with a 63x, 1.4 NA oil objective
- 542 using bidirectional scanning.

⁵⁴³ Data collection for the cluster analysis (*Figure 4*) results were performed on a Zeiss 980 laser ⁵⁴⁴ scanning confocal microscope set to use the Airvscan2 (ZEISS, Iena, Germany). Dorsal-mVenus and

⁵⁴⁵ MCP-mCherry were excited with argon ion laser lines wavelengths of 488 nm and 561 nm.

546 4.12 Image Analysis

Image processing and extraction of MS2 movies was performed in MATLAB using the custom pipeline described in (*Garcia et al., 2013*) and (*Lammers et al., 2020a*), which can be found in the public mRNADynamics Github repository. Transcription spots and nuclei were segemented with

the aid of the Trainable Weka Segmentation plugin for FIJI (*Witten et al., 2011*; *Arganda-Carreras et al., 2017*).

552 4.12.1 Bleaching quantification

⁵⁵³ Both the Dorsal-meGFP and Dorsal-mVenus curves in *Figure 3*B were quantified by generating

⁵⁵⁴ summed z-projections at each time point and calculating the fluorescence per pixel (summed fluo-⁵⁵⁵ rescence intensity in the 2D z-projection divided by the total number of pixels) for each time point.

⁵⁵⁵ rescence intensity in the 2D z-projection divided by the total number of pixels) for each time point.
 ⁵⁵⁶ All time points were then normalized to the first time point by dividing by the total fluorescence

⁵⁵⁷ intensity per pixel at t=0.

558 4.13 Single-molecule tracking

Single molecule imaging was performed on a Multimodal Optical Scope with Adaptive Imaging
 Correction (MOSAIC) at the Advanced Bioimaging Center (ABC) at UC Berkeley. The MOSAIC was in
 lattice light sheet imaging mode (*Chen et al., 2014*), without adaptive optics.

An initial snapshot of non-photoconverted Dorsal-mEos4a or Dorsal-Dendra2, excited by a 488 nm laser was taken to locate the nuclei in the embryo. Single-molecules were detected by first photoconverting a fraction of the Dorsal-mEos4a or Dorsal-Dendra2 molecules with a 405 nm laser and exciting the photoconverted molecules with a 560 nm laser with a 500 ms exposure time. Light sheet excitation was conducted with a Special Optics 0.65 NA, 3.74 mm working water dipping lens. Fluorescence emissions were detected with a Zeiss 1.0 NA water-dipping objective (2.2 mm working distance) and recorded approximately twice a second for the duration of the movie with a 2x Hamamatsu Orca Flash 4.0 v3 sCMOS camera.

Additional Lattice Light Sheet Microscopy (LLSM) (*Chen et al.*, 2014) experiments were per-570 formed at the Advanced Imaging Center at HHMI Janelia Research Campus. Embryos prepared 571 as described above were placed into the LLSM bath containing room temperature PBS. The full 572 details of the lattice light sheet microscope configuration are described previously (*Chen et al.*, 573 2014). A custom Special Optics 0.65 NA, 3.74 mm working distance water dipping objective was 574 used for excitation and a Nikon 1.1 NA, 2 mm working distance 25x water dipping objective (CFI 575 Apo LWD 25XW) was used for detection. A square lattice pattern (Inner NA: 0.44; Outer NA: 0.55) 576 was used for generating the lattice light sheet. Photoconversion was performed using a 405 nm 577 laser line while 488 nm and 560 nm laser lines were used for imaging. Emission was directed to 578 two Hamamatsu Orca Flash 4.0 sCMOS cameras (Dichroic: Semrock FF560-FDi01-25x36: Camera 1 570 Semrock BLP01-532R-25, Semrock NF03-488E-25, Semrock NF03-561E-25; Camera 2 – Semrock 580 BLP01-488r-25, Semrock FF01-520/35-25). The net system magnification is 63x for a pixel size of 581

104 nm. For all experiments, a field of view of 608 x 256 pixels was used.

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Custom scripts for the LLSM were developed for the photoconversion and single molecule imag-583 ing experiments. For both the Dorsal-mEos4a experiment and the H2B-mEos3.2 control experi-584 ment, an initial image of the histone channel was acquired (488 nm excitation, 100 ms exposure 585 time. 23 μ W: note that all powers were measured entering the back focal plane of the excitation 586 objective). For the Dorsal-mEos4a experiments, a 405 nm light sheet was scanned 10 μ m in 100 587 nm steps (101 images total) around the image plane (100 ms exposure time, 12.7 μ W). Due to the 588 higher labeling density of the H2B-mEos3.2, no initial photoconversion was necessary but instead 580 photobleaching was required to obtain an appropriate number of localizations per frame. To do 590 so, a 560 nm light sheet was scanned 5 μ m in 50 nm steps (101 images total) around the image 591 plane (100 ms exposure time, 748 μ W). This was repeated continuously for 30 iterations. All sub-592 sequent steps were identical for both the Dorsal-mEos4a experiments and H2B-mEos3.2 control 593 experiments. A single plane was imaged using excitation via a 560 nm light sheet. Separate exper-594 iments were performed with 3 different exposure times (50 ms, 100 ms, and 500 ms); a total of 595 2000, 1000, and 300 images were collected with excitation powers of 690 μ W, 455 μ W, and 153 μ W. 596 respectively. Due to the readout speed of the camera, the actual frame rate for each experiment 597 was 19.51 Hz, 9.88 Hz, and 2.00 Hz, respectively. After the single-plane, single-molecule imaging 598 was completed, a final image of the histone channel was collected (100 ms exposure time, 113 590 μ W) to account for any nuclear movement and assess changes in the stage of development. No 600 deskewing was necessary for the LLSM experiments as a single plane was imaged rather than a 3D 601 stack. 602

5 Supplementary Material

⁶⁰⁴ Supplementary Material, including figures and tables, can be found at the end of this document.

605 6 Acknowledgments

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622 **7** Author contributions

- 623 Conceptualization: MAT, HGG
- 624 Methodology: MAT, BM, NG, NL, HGG
- 625 Resources: MAT, GM, HGG
- 626 Investigation: MAT, BM, NG, NL, HGG
- 627 Visualization: MAT, HGG

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- 628 Funding acquisition: HGG
- 629 Project administration: HGG
- 630 Supervision: HGG
- 631 Writing original draft: MAT
- 632 Writing review & editing: MAT, HGG

8 Declaration of interests

⁶³⁴ The authors declare no competing interests.

9 Data and materials availability

636 All materials are available upon request. All data in the main text or supplementary materials are

₆₃₇ available upon request.

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Allele	CRISPR efficiency ^a	Homozyg. females produce pupae at 22°C ⁶	Homozyg. females produce pupae at 25°C ^b	Embryo hatch rate ^c	Rescues <i>dl</i> null allele ^d
<i>dl</i> (wild-type)	N/A	yes	yes	0.85	yes
dl-6G-mVenus-DsRed	unknown	yes	yes	0.75	yes
dl-6G-Dendra2-DsRed (line 1)	0.05	yes	no	0.55	*
dl-6G-Dendra2-DsRed (line 2)				0.60	—
dl-6G-meGFP-DsRed (line 1)	0.10	yes	no	0.05	—
dl-6G-meGFP-DsRed (line 2)				0.20	—
dl-6G-mEos4a-DsRed (line 1)	0.05	yes	no	0.15	—
dl-6G-mEos4a-DsRed (line 8)				0.20	—
dl-6G-mNeonGreen-DsRed (line 2)	0.06	yes	no	0.02	no
dl-6G-mNeonGreen-DsRed (line 6)				0.05	no
dl-LL-mNeonGreen-DsRed	0.60	yes	no	0.01	—
dl-6G-mNeonGreen-∆DsRed	N/A	yes	—	—	yes
dl-6G-mEos4b-DsRed	0.14	no	N/A	0.00	—
dl-6G-mEos3.2-DsRed	0.10	no	N/A	0.00	—
dl-6G-10GS-mEos3.2-DsRed	unknown	no	N/A	N/A	N/A
dl-6G-helix-mEos3.2-DsRed	unknown	no	N/A	N/A	N/A
dl-LL-mEos3.2-DsRed	unknown	no	N/A	N/A	N/A

Table 1. Summary of the *dl* **alleles characterized in this study.** For alleles with more than one entry, two unique CRISPR/Cas9 integrations were made into stable lines (labelled as e.g. "line 1" and "line 8") and characterized. Embryo hatch rate was quantified separately for each integration; all other metrics were combined between the two integrations of the same allele. Linker protein sequences: 6G (GGGGGG); LL (LongLinker; SGDSGVYKTRAQASNSAVDGTAGPGSTGSS); 6G-10GS (GGGGGGGGGGGGGGGGGGSGSGGS); 6G-helix (GGGGGGGMSKGEEL; MSKGEEL is the N-terminal helix from mVenus).

^{*a*} The number of DsRed+ adults divided by the total number of injected embryos that survived to adulthood. "Unsuccessful" indicates no injected embryos survived to adulthood.

^b females homozygous for the CRISPR allele (i.e. females deposit no wild-type Dorsal into the embryo) produce viable pupae.

^{*c*} Fraction of embryos laid by homozygous females that hatch after 36 hours.

^{*d*} One copy of the CRISPR allele complements (rescues function) a *dl* null allele. Either *dl*[1] or *dl*[4] were used.

* Not tested

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⁷⁹⁷ Supplementary Information

S1 Eliminating off-target CRISPR/Cas9 mutations does not improve viabil ity

We sought to remove any off-target mutations at other genomic loci by crossing the four new Dor-800 sal fusion lines with w; +; + flies for 6–8 generations, in a process called out-crossing or chromo-801 some cleaning (Methods Section 4.3). We then conducted an embryo hatch test using homozygous 802 females from these out-crossed lines, comparing the percentage of embryos laid after 36 hours to 803 the original CRISPR alleles. As shown in *Figure S1*, only one of the *dl-6G-meGFP-DsRed* lines saw an 804 increase in their hatch rates after outcrossing, but this improvement was only modest (from 5% to 805 8%) and did not improve embryo viability enough to enable maintenance of a stable population. 806 These results suggested that off-target CRISPR mutations were not a significant cause of lowered 807 viability in our new Dorsal fusion lines. 808

S2 Modifying the linker sequence does not improve viability

We also attempted to determine if the linker sequence was impacting the viability of our new fu-810 sion lines. When designing protein fusions, the length, flexibility, and composition of the linker 811 placed between the protein of interest and fluorescent protein can all be critical to maintaining en-812 dogenous activity and function (*Chen et al.*, 2013). We replaced the 6G linker (used by both *Reeves* 813 et al. (2012) and Alamos et al. (2023)) in the dl-6G-mNeonGreen-DsRed plasmid with another linker. 814 LL (literally, LongLinker; SGDSGVYKTRAQASNSAVDGTAGPGSTGSS, a kind gift of Michael Stadler). 815 However, only 1% of embryos from females homozygous for the modified linker dl-LL-mNeonGreen-816 DsRed allele hatched, and only after outcrossing (Figure S1). 817 Additionally, in an attempt to generate a mEos3.2 fusion allele that produces any viable progeny. 818 we replaced the 6G linker in the *dl-6G-mEos3.2-DsRed* plasmid with three other linkers: 6G-10GS 819 (GGGGGGGGGGGGSGSGS), LL, and 6G-helix (GGGGGGMSKGEEL, where MSKGEEL is the N-terminal 820 helix of mVenus). No viable progeny were produced by females homozygous for any of these three 821

alternative linker mEos3.2 alleles. Thus, the choice of linker did not have a stronger effect than the
 choice of fluoroscent protein.

⁸²⁴ S3 Quantification of nuclear Dorsal-mNeonGreen absolute concentration

⁸²⁵ We measured the absolute concentration of nuclear Dorsal-mNeonGreen fusion proteins in the ⁸²⁶ ventral nuclei of *DsRed* and $\Delta DsRed$ embryos using Raster Image Correlation Spectroscopy (RICS), ⁸²⁷ which can quantify the concentration of fluorescent protein molecules in live cells (*Digman et al.,* ⁸²⁸ 2005*a*,b; *Digman and Gratton, 2009; Schloop et al., 2024; Brown et al., 2008; Al Asafen et al., 2024*) ⁸²⁹ (*Section 4*). The nuclei on the ventral side of the embryos were imaged from nuclear cycle 10 until ⁸³⁰ gastrulation.

RICS analysis showed that the nuclear Dorsal-mNeonGreen concentration in the DsRed embryos 831 was lower than the concentration in the $\Delta DsRed$ across all nuclear cycles (*Figure S2*A). At the con-832 centration maxima in nuclear cycles 13 and 14. Dorsal-mNeonGreen concentration in DsRed em-833 bryos was only 36% and 47% of the concentrations measured in the nuclei of $\Delta DsRed$ embryos. 834 respectively (Figure S2B). The protein level is not reduced exactly to half as reported by our gPCR 835 measurements (Figure 2A). This discrepancy might result from errors in the estimation of concen-836 tration in RICS, due to variation in the mRNA degradation rate in the two lines, or due to differences 837 in translation efficiency of the dorsal mRNA. 838

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Figure S1. Embryo hatch test results before and after outcrossing. Hatch rates for embryos laid by females homozygous for the Dorsal fusion alleles, quantified as fraction of embryos that had hatched 36 hours after being laid. For the *dl-6G-Dendra2-DsRed2*, *dl-6G-meGFP-DsRed2*, *dl-6G-mEos4a-DsRed2*, and *dl-6G-mNeonGreen-DsRed2* alleles, homozygous females from original (top, dark bar in each pair) and outcrossed (bottom, light bar in each pair) versions of each fly line are compared. Embryo hatch rates for an untagged *dl* allele (from *yw* flies) served as a control.



Figure S2. Quantification of nuclear Dorsal-mNeonGreen absolute concentration. (A) Absolute Dorsal-mNeonGreen protein concentration, as quantified by RICS, in the ventral nuclei of DsRed (pink) and Δ DsRed (blue) embryos during nuclear cycles 10 to 14. Error bars are the standard error of the mean. **(B)** Ratio of absolute Dorsal-mNeonGreen protein concentration in Δ DsRed to DsRed embryos in ventral nuclei. DsRed: *dl-6G-mNeonGreen-DsRed* allele; Δ DsRed: *dl-6G-mNeonGreen-\DeltaDsRed* allele.

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Figure S3. Gene expression of Dorsal-activated genes in DsRed and Δ **DsRed embryos. (A)** Gene expression of two Dorsal-activated genes, *snail* (*sna*; green) and *short gastrulation* (*sog*; red), in the cross-section of a representative *dl-6G-mNeonGreen-\DeltaDsRed* embryo, as measured by fluorescence *in situ* hybridized (FISH). 24 embryos were imaged in total. *sna* is a Type I Dorsal-regulated gene, which is activated only by the highest levels of Dorsal on the ventral side of the embryo. *short gastrulation* (*sog*) is a Type III Dorsal-regulated gene, which is activated by even the lower levels of Dorsal present on the lateral sides of the embryo (*Reeves and Stathopoulos, 2009*) and repressed by high levels of Snail protein on the ventral side of the embryo. These expression patterns are similar to those seen in wild-type embryos (*Leptin, 1991*; *Francois et al., 1994*; *Srinivasan et al., 2002*). (**B**) Gene expression of the two Dorsal-activated genes, *sna* (green) and *sog* (red), in the cross-section of *dl-6G-mNeonGreen-DsRed* embryos, as measured by fluorescence *in situ* hybridized (FISH). 44 embryos were imaged in total. Representative images are shown for each of the three distinct types of gene expression patterns observed: (i) 11% (5/44) of embryos exhibit normal expression of both *sna* and *sog*; (ii) 66% (29/44) of embryos exhibit a narrow *sna* pattern and, in turn, an extended *sog* pattern; and (iii) 23% (10/44) of embryos exhibit an absent *sna* pattern and, in turn, an extended *sog* pattern.

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Figure S4. Single molecule detection of Dorsal-mEos4a and Dorsal-Dendra2 fusion proteins in live embryos. Single-molecule detection in an embryo expressing (A-D) Dorsal-mEos4a and (E-H) Dorsal-Dendra2. (A) Snapshot of non-photoconverted Dorsal-mEos4a excited by a 488 nm laser line to show the location of five ventral nuclei with high nuclear Dorsal levels. Nuclei 1 and 2 are labeled with white text. (B) Movie stills showing a series of single-molecule detections of a photoconverted portion of Dorsal-mEos4a molecules, which were photoconverted by a 405 nm laser to be excitable by a 560 nm laser. Images were taken with a 500 ms exposure of 560 nm light approximately twice a second. (C-D) Image series for the two nuclei labeled in (A), nucleus 1 (top) and nucleus 2 (bottom), showing (D) single-molecule detections of photoconverted Dorsal-mEos4a specifically within (C) the boundaries of each nucleus. (E-H) Single-molecule detection in an embryo expressing Dorsal-Dendra2. (E) Snapshot of non-photoconverted Dorsal-Dendra2 excited by a 488 nm laser line to show the location of five ventral nuclei with high nuclear Dorsal levels. Nucleus 3 is labeled with white text. (F) Movie stills showing a series of single-molecule detections of a photoconverted portion of Dorsal-Dendra2 molecules, which were photoconverted by a 405 nm laser to be excitable by a 560 nm laser. Images were taken with a 500 ms exposure of 560 nm light approximately twice a second. (G-H) Image series for the nucleus 3, labeled in (E), showing (H) single-molecule detections of photoconverted Dorsal-Dendra2 specifically within (G) the boundaries of the nucleus.

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Transgenic Fly Lines				
Genotype	Source			
w[1118] ; + ; PBacy[+mDint2] GFP[E.3xP3]=vas-Cas9VK00027	BDSC #51324			
w[1118]	BestGene, Inc.			
уw	BestGene, Inc.			
уw ; Sp/CyO ; +	lab stock			
yw / w ; Dorsal-6G-mVenus-DsRed ; +	Alamos et al. (2023)			
yw / w ; Dorsal-6G-meGFP-DsRed / CyO ; +	current study			
yw / w ; Dorsal-6G-mNeonGreen-DsRed / CyO ; +	current study			
yw / w ; Dorsal-6G-Dendra2-DsRed / CyO ; +	current study			
уw / w ; Dorsal-6G-mEos4a-DsRed / СуО ; +	current study			
уw / w ; Dorsal-6G-mEos4b-DsRed / СуО ; +	current study			
yw / w ; Dorsal-6G-mEos3.2-DsRed / CyO ; +	current study			
yw / w ; Dorsal-10GS-mEos3.2-DsRed / CyO ; +	current study			
yw / w ; Dorsal-6G-helix-mEos3.2-DsRed / CyO ; +	current study			
yw / w ; Dorsal-LongLinker-mEos3.2-DsRed / CyO ; +	current study			
yw / w ; Dorsal-LongLinker-mNeonGreen-DsRed / CyO ; +	current study			
yw; Dorsal-mVenus-DsRed, pNos-MCP-mCherry; Dorsal-mVenus,	Alamos et al. (2023)			
pNos-MCP-mCherry, His2Av-iRFP				
уw; eNosx2-MCP-mCherry / СуО ; +	current study			
yw; Dorsal-meGFP-DsRed, eNosx2-MCP-mCherry / CyO ; +	current study			
yw; Dorsal-mNeonGreen-DsRed, eNosx2-MCP-mCherry / CyO ; +	current study			
w; snaBAC-MS2; +	Bothma et al. (2015)			
yw ; hbP2P-MS2 ; +	Garcia et al. (2013)			

Table S1. List of fly lines used in the current study.

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Plasmids				
Name	Source			
pU6-DorsalgRNA1	Alamos et al. (2023)			
dl-6G-mVenus-DsRed	Alamos et al. (2023)			
dl-6G-meGFP-DsRed	current study			
dl-6G-mNeonGreen-DsRed	current study			
dl-6G-Dendra2-DsRed	current study			
dl-6G-mEos4a-DsRed	current study			
dl-6G-mEos4b-DsRed	current study			
dl-6G-mEos3.2-DsRed	current study			
dl-10GS-mEos3.2-DsRed	current study			
dl-6G-helix-mEos3.2-DsRed	current study			
dl-LongLinker-mEos3.2-DsRed	current study			
dl-LongLinker-mNeonGreen-DsRed	current study			
snailBAC/MS2-yellow (snaBAC)	Bothma et al. (2015)			
pIB-hbP2P-24xMS2v5-lacZ-tub3'UTR	Garcia et al. (2013)			

Table S2. List of plasmids described in this study. Full sequences for all plasmids introduced in the current study can be accessed through a Benchling repository at https://benchling.com/garcialab/f_/THClp5A3-dorsal-fusions-manuscript/









A





A Dorsal-mEos4a + H2B-GFP 488 nm excitation

1

B Dorsal-mEos4a 560 nm excitation





Α





0 min

2 min

6 min



Е