# The Spatio-Temporal Control of Zygotic Genome Activation

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## SUMMARY

- 8 One of the earliest and most significant events in embryonic development is zygotic
- 9 genome activation (ZGA). In several species, bulk transcription begins at the mid-
- 10 blastula transition (MBT) when, after a certain number of cleavages, the embryo attains
- 11 a particular nuclear-to-cytoplasmic (N/C) ratio, maternal repressors become sufficiently
- diluted, and the cell cycle slows down. Here we resolve the frog ZGA in time and space
- 13 by profiling RNA polymerase II (RNAPII) engagement and its transcriptional readout. We
- 14 detect a gradual increase in both the quantity and the length of RNAPII elongation before
- 15 the MBT, revealing that >1,000 zygotic genes disregard the N/C timer for their activation,
- 16 and that the sizes of newly transcribed genes are not necessarily constrained by cell
- 17 cycle duration. We also find that Wnt, Nodal and BMP signaling together generate most
- 18 of the spatio-temporal dynamics of regional ZGA, directing the formation of orthogonal
- 19 body axes and proportionate germ layers.

## 20 INTRODUCTION

- 21 The genomes of multicellular organisms are transcriptionally silent at the time of fertilisation,
- and the events of early development, including zygotic (also known as embryonic) genome
- activation (ZGA), are directed by maternal gene products (De Iaco et al., 2019; Eckersley-
- Maslin et al., 2019; Gentsch et al., 2018b; Lee et al., 2013; Leichsenring et al., 2013; Liang et
- al., 2008). The number of cell cycles after which ZGA becomes essential for development (at
- which embryos arrest if transcription is inhibited) is highly reproducible within each species. In
- 27 the zebrafish, the frog *Xenopus* and the fruit fly *Drosophila*, this occurs after 10, 12 and 13 cell
- 28 cycles, respectively, at the so-called mid-blastula transition (MBT) (Blythe and Wieschaus,
- 29 2015; Gentsch et al., 2018b; Kane and Kimmel, 1993; Newport and Kirschner, 1982a). Early
- 30 development in these species occurs with no gain in cytoplasmic volume, and studies in
- 31 Xenopus suggest that ZGA is triggered at a particular nuclear-to-cytoplasmic (N/C) ratio, when
- 32 the increasing amount of nuclear DNA titrates out maternally deposited repressors (Newport

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and Kirschner, 1982b). Slower-developing mammalian embryos show major waves of RNA polymerase II (RNAPII)-mediated transcription as early as the 2-cell stage in mice (Bolton et al., 1984; Hamatani et al., 2004) and 4- to 8-cell stage in humans (Braude et al., 1988; Vanessa et al., 2011). This occurs days before the formation of the blastocyst, which, like the blastula, contains the pluripotent cells that form the embryo proper. In Xenopus, ZGA is associated with changes in cell behaviour after the MBT. First, rapid and nearly synchronous cell cleavages give way to longer and asynchronous cell divisions (Anderson et al., 2017; Newport and Kirschner, 1982a). Second, embryonic cells acquire the ability to respond to inductive signaling (Gentsch et al., 2018b), causing them to become motile, to establish dorso-ventral patterning, and to contribute to one or two of the three germ layers (endoderm, mesoderm and ectoderm). These germ layers emerge first during gastrulation and are the primordia of all organs. Third, embryos show accelerated degradation of maternal RNA, and fourth, cells gain apoptotic (Stack and Newport, 1997) and immunogenic (Gentsch et al., 2018a) capabilities. While large-scale ZGA occurs at the MBT, some genes escape the repressive environment of the early embryo, and nascent transcripts can be detected in Xenopus during rapid cleavage stages. For example, primary microRNA transcripts of the polycistronic MIR-427 gene (Lund et al., 2009) are detectable in X. tropicalis after just three cell divisions (Owens et al., 2016). MIR-427, like its zebrafish equivalent MIR-430, is activated at early stages by the synergistic and pioneering activities of maternal members of the SoxB1 and Pou5F (Oct4) transcription factor (TF) families (Gentsch et al., 2018b; Heyn et al., 2014; Lee et al., 2013). These core pluripotency TFs, represented by Sox3 and Pou5f3 in Xenopus, are characterized by ubiquitous and high translation frequencies in pre-MBT embryos (Gentsch et al., 2018b; Lee et al., 2013). Zygotic transcription of the Nodal-encoding genes nodal3/5/6, and of the homeobox genes siamois1/2, is initiated by nuclear β-catenin as early as the 32-cell stage (Owens et al., 2016; Skirkanich et al., 2011; Yang et al., 2002). While miR-427 (and miR-430 in zebrafish) contributes to the clearance of maternal RNA (Giraldez et al., 2006; Lund et al., 2009), nodal and siamois gene initiate the formation of the germ layers and body axes (Agius et al., 2000; Lemaire et al., 1995). All these genes, and other early-activated genes in *Drosophila*, *Xenopus* and zebrafish, have coding sequences of <1 kb and either no introns or just a few (Heyn et al., 2014). It has been suggested that the early rapid cell cycles cause DNA replication machinery to interfere with the transcription of larger genes (Shermoen and O'Farrell, 1991), a suggestion supported, to date, by the profiling of nascent transcripts (Heyn et al., 2014). We note, however, that the detection and temporal resolution of de novo transcription can be particularly challenging for genes that have both maternal and zygotic transcripts. Here we use the continuous occupancy of RNAPII along gene bodies as a method to record ZGA. In contrast to transcript profiling techniques, this method (1) directly determines the

activity of every gene; (2) is independent of metabolic labeling (Heyn et al., 2014) and of any gene feature such as introns (Lee et al., 2013), single nucleotide polymorphisms (Harvey et al., 2013; Lott et al., 2011) or transcript half-lives; and (3) circumvents difficulties in detecting nascent transcripts in a large pool of maternal transcripts. By combining these data with the profiling of the transcriptome along the primary body axes (Blitz et al., 2017), we resolve ZGA in time and space for wild-type and various loss-of-function embryos. We provide evidence that runs counter to our original understanding of the cell cycle or of the N/C ratio in constraining gene expression before MBT. And finally, we show how signaling initiates and coordinates spatio-temporal ZGA in the *Xenopus* embryo.

## **RESULTS**

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# **RNAPII Profiling Reveals Exponential ZGA before MBT**

In an effort to resolve the progression of ZGA, we profiled chromatin for RNAPII engagement on hand-sorted X. tropicalis embryos over six developmental stages from the 32-cell to the late gastrula stage (Figure 1A,B). RNAPII was localized on the genome by chromatin immunoprecipitation followed by deep sequencing (ChIP-Seq). We complemented RNAPII profiling with high time-resolution transcriptomics (Owens et al., 2016) counting both exonic and intronic RNA at 30-min intervals from fertilization to the late gastrula stage (Figures 1A,B and S1A,B). For both maternal and zygotic genes, the detection threshold was set to ≥3 transcripts per million (TPM) averaged over any 1-h window during this developmental period to avoid genes with general low-level expression. This restricted the analysis to 13,042 genes (Figure 1B). These genes were considered active when we detected simultaneously RNAPII enrichment along their full length (see Transparent Methods) as well as the presence of the corresponding transcripts. In doing so, we used a low threshold of ≥0.1 TPM so as not to miss the onset of gene transcription. RNAPII-guided ZGA profiling was verified in part by active posttranslational histone marks (Hontelez et al., 2015) and by differential expression methods aiming at detecting nascent transcripts. Thus, zygotic transcript depletion (by blocking RNAPII elongation with α-amanitin) (Gentsch et al., 2018b) or enrichment (by selecting 4-thiouridine [4sU] tagged transcripts at the MBT and the mid-gastrula stage) showed substantial overlaps and positive correlations with RNAPII-covered genes (Figures 1A,B and S1C,D and Tables S1 and S2).

This analysis revealed an exponential ZGA before the MBT with 27, 144 and 1,044 active genes after 5 (32-cell, ~2.5 hpf), 7 (128-cell, ~3 hpf) and 10 (1,024-cell, ~4 hpf) cell cycles, respectively. Gene activation reached its peak at the MBT (~4.5 hpf), with 1,854 newly-activated genes, before dropping to 724 genes at the early-to-mid gastrula stage (~7.5 hpf) and increasing again to 1,214 genes towards the end of gastrulation (~10 hpf) (Figures 1B,C and S1E and Table S2). The dramatic increase in transcriptional activity that occurs in the 1.5 hours between the 128-cell stage and the MBT can be illustrated by Hilbert curves (Figure 1C), which

provide a genome-wide overview of RNAPII enrichment by folding chromosomes into two-dimensional space-filling and position-preserving plots (Gu et al., 2016). While most zygotic genes remain active beyond the mid-gastrula stage, 197 (including *siamois2* [*sia2*], *nodal5* and *znf470*) of the 4,836 zygotic genes (~4%) are deactivated within ~6 h of development (Figures 1C and S1F,G). Slightly less than one third of the activated genes were differentially expressed along either or both of the animal-vegetal and dorso-ventral axes (Figures 1D and S1F).

The temporal order of enriched biological processes supported by ZGA matched the regulatory flow of gene expression, starting with nucleosome assembly, nucleic acid synthesis, mRNA metabolism and production, post-translational modification and degradation of proteins (Figure 1E). The earliest transcriptional engagement, beginning at the 32 to 128-cell stages, was detected in gene clusters of tens to hundreds of kilobases (Figures 1A,C and S1H-J). These clusters featured close relatives of the same genes, some of which are critical to Nodal signaling (Nodal ligands), the formation of the Spemann organiser (Siamois homeobox transcription factors), nucleosome assembly (histones), mRNA decay (MIR-427), and ongoing gene regulation (zinc finger [ZF] transcription factors with on average 10 Cys2-His2 [C2H2] domains; Figure S1J). These earliest activated genes were shorter and encoded smaller proteins than those within the maternal pool or those that are activated post-MBT (Figures 1F and S1K). The non-coding features that contributed most to the differences in length were the 3' UTRs and introns (Figure S1K).

We noted that the shorter zygotic genes observed before the MBT did not correlate strictly with the time constraints imposed by short cell cycles. We detected increasing and wider spread of de novo recruitment of RNAPII before the MBT, when cleavages continue to occur at rapid and near-constant pace (Figures 1F and S1K). During this period, the median length of activated genes (and their coding sequences) increases from ~0.9 kb (~0.4 kb) to ~5.9 kb (~0.9 kb). However, it was not until after the MBT that the overall architecture of zygotic and maternal genes became indistinguishable (Figures 1F and S1K). Temporal comparison of RNAPII engagement and total RNA profiling suggested that the zygotic contribution to the transcriptome (as calculated by the number of zygotic genes divided by the number of genes with ≥0.1 TPM maternal transcripts averaged over the first hour post-fertilisation when the entire zygotic genome is still transcriptionally inactive) rose within seven cell cycles from ~0.2% at the 32-cell stage to ~22% at the MBT (Figure 1G). Further maternal degradation and more moderate transcriptional engagement extended the zygotic contribution to about one third of the transcriptome by the late gastrula stage. Maternal transcripts (≥0.1 TPM, see above) were detected for ~67% of newly activated genes (18 out of 27 genes) at the 32-cell stage, ~85% (99/117) at the 128-cell stage, ~87% (780/900) at the 1,024-cell stage, ~95% (1,754/1,854) at the MBT, ~89% (644/724) at the mid-gastrula stage and ~90% (1,094/1,214) at the late gastrula stage (Table S2). Altogether ~91% (4,389/4,836) of newly activated genes have ≥0.1 TPM maternal contribution.

Wnt, Nodal and BMP Signals Are Key Drivers of Regional ZGA

We next sought to investigate the single and combined effects of different inductive signals on the spatio-temporal dynamics of ZGA. The early vertebrate embryo employs canonical Wnt, Nodal and BMP signals and their key transcriptional effectors β-catenin, Smad2 and Smad1, respectively, to establish the primary body axes and the three germ layers (reviewed by Arnold and Robertson, 2009 and Kimelman, 2006). In *Xenopus*, β-catenin first translocates to the nuclei of dorsal blastomeres at the 32-cell stage (Larabell et al., 1997; Schneider et al., 1996) (Figure 2A). After the MBT, zygotic Wnt8a causes more nuclear β-catenin to accumulate around the forming blastopore lip (Christian and Moon, 1993; Schohl and Fagotto, 2002). The nuclear translocation of Smad1 and Smad2 is triggered around the MBT by various BMP and Nodal ligands. Nuclear Smad1 is primarily detected on the ventral side and the blastocoel roof of the embryo while nuclear Smad2 is detected within the vegetal hemisphere (VH) and the marginal zone (MZ) (Faure et al., 2000; Schohl and Fagotto, 2002) (Figure 2A).

In an effort to inhibit canonical Wnt signaling, we injected into the X. tropicalis zygote a previously validated antisense morpholino oligonucleotide (MO) which interferes with β-catenin protein synthesis by annealing to the translation start codon (Heasman et al., 2000). Nodal and BMP signals were selectively blocked by incubating dejellied embryos from the 8-cell stage in the cell-permeable inhibitors SB431542 (Ho et al., 2006; Inman et al., 2002) and LDN193189 (Cuny et al., 2008; Young et al., 2017), respectively. The morphological phenotypes of these single loss-of-function (LOF) treatments were consistent with previous observations and ranged from impaired axial elongation causing the loss of tail structures (BMP, Reversade et al., 2005) to severe gastrulation defects (Wnt, Heasman et al., 2000, and Nodal, Ho et al., 2006) as shown in Figure 2B. Briefly, Nodal LOF impaired blastopore lip formation and bulk tissue movements of gastrulation (bullet points in Figure 2B). However, it did not preclude subsequent elongation of the antero-posterior axis. By contrast, Wnt LOF embryos underwent gastrulation (albeit delayed and more circumferentially rather than in a dorsal to ventral wave), but failed to form an antero-posterior axis, with both head and tail being absent. With respect to the joint effects of Wnt, Nodal and BMP signaling, most dual and triple LOFs combined their individual morphological defects such that, for example, Wnt/Nodal LOF resulted in the complete loss of gastrulation and axial elongation. By contrast, Wnt/BMP LOF produced defects such as nonfusing neural folds (arrowheads in Figure 2B), structures that were either absent in Wnt LOF embryos or normal in BMP LOF embryos.

Changes to ZGA caused by the single or combined LOF of Wnt, Nodal and/or BMP were then determined at the late blastula stage on a transcriptome-wide scale using deep RNA sequencing (RNA-Seq). Analysis was limited to the 3,315 zygotic genes for which spatio-temporal expression data is available (Blitz et al., 2017; Owens et al., 2016) and where reduced expression ( $\geq$ 50% loss of exonic and/or intronic transcript counts, FDR  $\leq$ 10%) could be detected in  $\alpha$ -amanitin-injected embryos (Figure 2C and Table S3) (Gentsch et al., 2018b).  $\alpha$ -Amanitin-mediated inhibition of RNAPII elongation impedes the morphogenetic tissue

movements of gastrulation and ultimately leads to early embryonic death (Gentsch et al., 2018b). Spatial gene expression patterns were inferred from experiments comparing the transcriptomes of embryos dissected along their animal-vegetal and dorso-ventral axes (Blitz et al., 2017); we did not include the left-right axis because there were no significant differences in gene expression across this axis at the gastrula stage (Blitz et al., 2017). The signal-mediated transcriptional effects (1.5-fold change from control RNA level) on zygotic genes, 86% (2840/3315) of which have ≥0.1 TPM maternal contribution, ranged from ~1.5% (~1.3% down and ~0.2% up) to ~26% (~19% down and 7% up) for single BMP LOF and triple Wnt/Nodal/BMP LOFs, respectively (Figure 2C). As expected, the transcript levels of genes whose expression is solely zygotic were more strongly affected than those of zygotic genes with maternally contributed transcripts (Figures 2C and 4A). The extent of ZGA mis-regulation largely reflected the severity of the resulting morphological phenotypes at the late gastrula and the mid-tailbud stage (Figure 2B,C).

In comparison, the LOFs of critical maternal TFs like Pou5f3/Sox3 or VeqT (Gentsch et al., 2018b) caused the mis-regulation of 61% (~24% down and ~37% up) and 13% (~6% down and ~7% up) of zygotic genes, respectively. The LOFs of four zygotic T-box TFs (zVegT, Eomes, Tbxt and Tbxt.2), all of which require Nodal signaling for their expression, caused slight misregulation in 19% (~9% down and ~10% up) of the zygotic genes as detected over three consecutive developmental time points during gastrulation (Table S3). Among the ZGAenriched biological functions (Figure 1D), Wnt, Nodal and BMP signals, like maternal Pou5f3/Sox3 and VeqT, strongly affected zygotic genes associated with cell migration, gastrulation, dorso-ventral and antero-posterior body axis formation and regionalization (Figure 2D). Impaired tissue movements during gastrulation, as observed in various LOFs (Figure 2B and Movie S1), was prefigured by a strong enrichment for cell migration-associated genes. The genes suppressed or unaffected by the selected signals and maternal TFs were enriched for the ZGA-critical biological processes of mRNA metabolism and translation. For instance, the transient activation of the entire zinc finger cluster (Figure S2A) was not affected by any tested LOF. Because family members are frequently cross-regulated, and the MBT-staged chromatin contains many Krüppel-like zinc finger 'footprints' (Gentsch et al., 2018b), it is conceivable that the unaffected, tissue-nonspecific part of ZGA is regulated by maternal zinc finger TFs. This vertebrate gene regulatory branch may be more ancient than that of Pou5F/SoxB1 as zinc finger TFs like Zelda are also key to ZGA of the invertebrate Drosophila (Liang et al., 2008).

Next, signal-dependent ZGA was resolved in time and space based on (i) the profiling of RNAPII-engaged chromatin from the 32-cell stage to the MBT and (ii) known gene expression patterns along the animal-vegetal and dorso-ventral axes (Blitz et al., 2017) (Figures 2E and S2B-F). In line with the nuclear translocation of their signal mediators (Figure 2A), Wnt, Nodal and BMP proved to be required for gene activation in different spatio-temporal domains of the early embryo:  $\beta$ -catenin was needed for ~87% and ~46% of genes preferentially expressed on the dorsal side and in the VH/MZ, respectively. Some of its target genes like *nodal3.1* and *sia2* 

were already active by the 32-cell stage (Figures 1C and 2E). Upon Wnt LOF, the early transcriptional down-regulation was followed by the mis-regulation of opposing cell fate

specifiers: the upregulation of ventral genes (e.g., id2, szl) and the downregulation of dorsal

genes (e.g., chrd, otx2). These observations suggest that β-catenin protects dorsal cells from

ventralization (Figures 2E and S2C).

Along similar lines, Nodal LOF embryos predominantly displayed a down-regulation of dorsal

(~63%) and VH/MZ-specific (~73%) genes, although there was no effect of Nodal LOF on the

earliest-activated genes at the 32-cell stage or on opposing cell fate regulators (Figure S2A,B).

Among the first genes to be sensitive to Nodal LOF was the MZ-specific FGF ligand fgf20,

activated by the 128-cell stage (Figures 1C and S2A,B). By contrast, BMP LOF caused a

decrease in ventrally-expressed gene expression (~45%) from the 1,024-cell stage onwards

235 (Figure S2A,C).

respectively).

As a comparison, the ubiquitous expression of maternal Pou5f3/Sox3 was required for transcription in all spatio-temporal domains, including, for example, the uniform expression of miR-427 (Figure S2A,D). The requirement for Pou5f3/Sox3 was more marked, however, for region-specific genes, in particular those expressed within animal- (~55%) and ventral-specific (~67%) domains (Figure S2A,D). The maternal TF VegT promoted vegetal identity by activating ~40% of genes transcriptionally enriched within its own expression domain, the vegetal hemisphere, while suppressing genes that are expressed in the animal hemisphere like *foxi4.2*. The requirement for VegT was similar in ventral- and dorsal-specific genes (~31% and ~30%

## Wnt/BMP Synergy Enables Uniform ZGA Across the Dorso-Ventral Axis

The relationships between inductive signals with respect to spatial aspects of ZGA was explored by comparing zygotic transcriptomes in single and double LOF experiments. Interestingly, while simultaneous loss of both Nodal and BMP function, or both Nodal and Wnt, caused additive effects on gene expression compared to the single LOFs (Figures 3E and S3A-N), simultaneous loss of both BMP and Wnt signaling showed a more synergistic effect (Figure 3A,F). These observations are consistent with the morphological consequences of single and double LOFs (Figure 2B). Single LOF experiments revealed very little overlap between Wnt and BMP gene targets (Figure 3B), a result consistent with their domains of activity which are, initially, at opposite ends of the dorso-ventral axis. However, dual Wnt/BMP inhibition increased the number of downregulated genes by 292, a rise of ~118% and ~664% with respect to individual Wnt- and BMP-dependent genes respectively (Figure 3B). Interestingly, this synergy affected 166 Nodal-dependent genes, most of which had uniform expression levels across the dorso-ventral axis (Figure 3C,D,F-H). Thus, spatially-restricted Wnt, BMP and Nodal signals act together to establish dorso-ventral expression uniformity of genes such as *tbxt* and *eomes* (Figure 3I).

Overall, the loss of canonical Wnt, Nodal and/or BMP signaling caused the misregulation of ~39% (~22.1% down, ~2.1% down/up and ~14.4% up) of genes activated at ZGA (Figure 4A). These signals were required for most regional ZGA on the dorsal side (~89%) and in the VH and MZ (~82%). Notably, their input affected virtually all genes (~98%, 56/57) with enriched expression in the dorso-vegetal/MZ quadrant (Figure 4B,C). Thus, Wnt, Nodal and BMP substantially contribute to regional ZGA in most anatomical domains of the early gastrula embryo with the exception of animally enriched transcription (~19%). Animal- and ventral-specific gene expression relies strongly on both activation by ubiquitous maternal TFs (e.g. Pou5f3/Sox3) and on repression by signals (Figure 4B,D) and other maternal TFs (e.g. VegT) on the opposite side (Figure S2F).

#### DISCUSSION

Our study provides two major insights into the mechanisms by which ZGA is initiated in time and space in *Xenopus tropicalis*. The first concerns the temporal aspects, where we find that RNAPII can be detected across gene bodies well before the MBT, during the period when rapid synchronous cell divisions divide the zygote into 4,096 blastomeres (Figure 4E). The average length of genes covered by RNAPII grows during this time, from ~1 kb at the 32-cell stage to ~6 kb at the 1,024-cell stage; these figures compare with an average size of ~16 kb for maternally-expressed genes and for genes expressed after the MBT. Recent long-read sequencing of the zebrafish transcriptome at pre-MBT stages identified transcripts as long as 8 kb spanning multiple pri-miR-430 elements (Nudelman et al., 2018). Furthermore, RNAPII elongation in pre-MBT *Drosophila* embryos occurred at rates of 2.4 to 3.0 kb/min (Chen et al., 2013; Fukaya et al., 2017).

We do not know why RNAPII, despite its high abundance and its ability to promote rapid elongation, is restricted at early stages from transcribing more genes and longer genes. It may, perhaps, be a consequence of the gradual nature of the chromatin remodeling that occurs during these stages, from the accessibility of *cis*-regulatory elements (Gentsch et al., 2018b; Liu et al., 2018; Lu et al., 2016; Wu et al., 2016) to the spatial organisation of an initially unstructured or highly variable chromatin landscape (Du et al., 2017; Flyamer et al., 2017; Hug et al., 2017; Kaaij et al., 2018; Ke et al., 2017). Whatever the reason, the increase of elongated RNAPII engagement between the 128-cell stage and the MBT indicates that a significant component of ZGA disregards the N/C ratio which was originally thought to underlie the onset of transcription at the MBT (Newport and Kirschner, 1982b). Similar conclusions have been drawn from profiling the zygotic transcriptome of haploid *Drosophila* (Lu et al., 2009) and cell cycle-arrested zebrafish (Chan et al., 2018). Thus, it is becoming clear from work in flies, fish and frogs that ZGA starts before the MBT and accelerates thereafter (Ali-Murthy et al., 2013; Collart et al., 2014; Mathavan et al., 2005; Owens et al., 2017; Tan et al., 2013), reaching a peak at the MBT (reviewed by Jukam et al., 2017 and Langley et al., 2014).

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These observations notwithstanding, it remains possible that cell cycles do contribute to the temporal progression of ZGA and the exponential increase in the number of activated genes before the MBT. In particular, cell cycles may accelerate chromatin remodeling by displacing suppressors in mitotic chromatin and providing unique access to TFs (Halley-Stott et al., 2014) and structural proteins of high-order chromatin (Ke et al., 2017). For example, maternal core histones have been shown to prevent premature ZGA by competing with specific TFs (Joseph et al., 2017). In addition to the small sizes of the earliest activated genes, we observed that most of these genes, which have no or few introns, code for groups of related factors like histones or zinc finger TFs, and that they appear as clusters spanning up to several hundred kilobases. This is in line with previous findings of the earliest active multicopy and intron-poor genes like miR-427 and nodal5/6 in Xenopus embryos (Collart et al., 2014; Lund et al., 2009; Owens et al., 2016; Skirkanich et al., 2011; Takahashi et al., 2006; Yanai et al., 2011; Yang et al. 2002) and miR-430 in zebrafish and Medaka fish (Giraldez et al., 2005; Heyn et al., 2014; Tani et al., 2010). The number and spatial proximity of clustered genes enhances transcriptional output by allowing the sharing of multiple cis-regulatory elements (arranged as super-enhancers) (Whyte et al., 2013) and by fortifying transcriptional condensates of TFs, coactivators and RNAPII (Boija et al., 2018; Cho et al., 2018; Chong et al., 2018; Sabari et al., 2018; Shrinivas et al., 2018). Overall, based on enriched gene functions, we discovered that ZGA exerts a temporal control of gene expression from nucleosome remodeling before the MBT to protein degradation after the MBT. Our second insight concerns spatial ZGA, and the observation that we can assign a large proportion of spatio-temporal ZGA to key signaling pathways (reviewed by Arnold and Robertson, 2009 and Kimelman, 2006). Canonical Wnt, Nodal and BMP signaling govern regional ZGA in line with the nuclear translocation of their signal mediators (Faure et al., 2000; Larabell et al., 1997; Schohl and Fagotto, 2002). Thus, Nodal signaling predominantly affects transcription within the vegetal hemisphere and marginal zone, while Wnt and BMP initiate transcription in dorsal and ventral regions, respectively. The timing of regional ZGA is defined by the sequential translocation of signal mediators such that nuclear β-catenin directs regional ZGA at the 32-cell stage, followed by nuclear Smad2 at the 128-cell stage and Smad1 at the 1,024-cell stage. While Smad2-mediated signal transduction depends on the zygotic transcription of its six Nodal ligands (Faure et al., 2000; Gentsch et al., 2018b; Jones et al., 1995; Yang et al., 2002), canonical Wnt and BMP signaling are initiated by the maternally inherited ligands Wnt11 and BMP2/4/7, respectively (Faure et al., 2000; Heasman, 2006; Tao et al., 2005). We also show that the synergy of opposing signals of the Wnt and BMP pathway affects many Nodal-dependent genes with uniform expression along the dorso-ventral axis such as eomes and Brachyury (tbxt). It is not yet clear whether Wnt/BMP synergy arises from joint chromatin

engagement or from mutual or post-translational interactions. For instance, Wnt8a signal can enhance BMP transcriptional readouts by inhibiting the phosphorylation of GSK3, which normally targets Smad1 for degradation (Fuentealba et al., 2007). However, the analysis of *Brachyury* gene regulation in zebrafish suggests that Wnt and BMP can be integrated at a single cis-regulatory DNA element and together with a separate Nodal-responsive DNA element they can establish uniform dorso-ventral expression (Harvey et al., 2010). This is further corroborated by our analysis of genome-wide chromatin engagement (Gentsch et al., 2018b): the canonical DNA recognition motif for the Wnt-associated basic helix-span-helix (bHSH) TF AP-2 was more enriched at Smad1 than Smad2 binding sites (Figure S3O).

We therefore propose that Wnt, BMP and Nodal signal mediators are critical to regional ZGA and that they balance initially opposing cell fate commitments. However, we have previously shown that signal integration also relies on maternal pioneer TFs like Pou5f3 and Sox3 to make signal-responsive *cis*-regulatory elements accessible for signal mediator binding. For example, Nodal-induced transcription of the *Brachyury* gene depends on the pioneering roles of maternal

Pou5f3 and Sox3, and less on their transcriptional activities (Gentsch et al., 2018b).

Overall, we demonstrate that the temporal and spatial dynamics of regional ZGA are regulated

by the sequential and spatially restricted translocation of Wnt, Nodal and BMP signal mediators.

These events establish the formation of the primary body axes and germ layers of the embryo.

Temporal RNAPII profiling indicates that >1,000 genes of increasing length are activated before

MBT and that this substantial portion of ZGA is independent of both the classic N/C ratio and

of cell cycle lengthening.

## LIMITATIONS OF THE STUDY

We detected a dramatic increase in genome-wide recruitment of RNAPII over the cleavage stages during which the genome begins to be transcribed. We used within-sample normalization to scale developmental stage-specific RNAPII profiles. However, because of the large differences in total RNAPII enrichment between samples, chromatin spike-ins are now considered a more accurate method to normalize ChIP-Seq profiles across consecutive developmental stages (Chen et al., 2016). We combined separate whole-embryo determinations of RNAPII engagement and transcript levels to reveal the temporal dynamics of ZGA. This approach could be improved by profiling RNAPII-associated RNA to directly couple RNAPII elongation with transcript accumulation (e.g., Churchman and Weissman, 2011). In addition, the spatial resolution of ZGA, which is based on transcriptomics of dissected embryonic parts in our study, could be enhanced by various deep single-cell profiling and super-resolution imaging technologies. We show that most regional ZGA depends on Wnt, Nodal and BMP signals, but an important question remains: How are these signals integrated at the chromatin level to sustain RNAPII-mediated transcript elongation? In part, this could be

- investigated by targeted genome editing to increase our understanding of signal-responsive
- gene regulatory DNA.

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## **AUTHOR CONTRIBUTIONS**

- 383 Conceptualization, G.E.G.; Methodology, G.E.G.; Computational Code, G.E.G.; Formal
- Analysis, G.E.G. and N.D.L.O.; Investigation, G.E.G.; Writing Original Draft, G.E.G. and
- J.C.S.; Writing Review & Editing, G.E.G and J.C.S.; Funding Acquisition, J.C.S.

## 386 **DECLARATION OF INTERESTS**

The authors declare no competing interests.

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## **MAIN FIGURES**

610

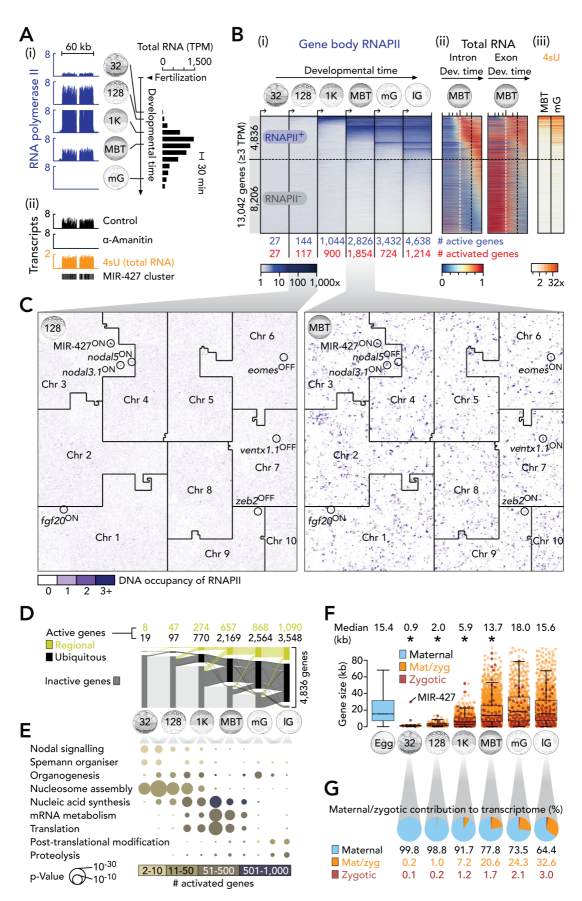


Figure 1. Dynamics and Architecture of ZGA in X. tropicalis

- 613 (A) (i) Genome-wide profiling of RNAPII and total RNA (Owens et al., 2016) to determine
- temporal ZGA dynamics. (ii) Complementary approach to detect transcriptionally active genes
- 615 by α-amanitin-induced loss and 4sU-enrichment of nascent (zygotic) transcripts.
- (B) Progression of ZGA from the 32-cell to the late gastrula stage based on (i) whole gene body
- 617 (full-length) occupancy of RNAPII (i.e., RNAPII was enriched across entire gene bodies; see
- Transparent Methods). Co-aligned: (ii) High time-resolution of total RNA, separated by intron-
- and exon-derived signals, from fertilization to the late gastrula stage, and (iii) enrichments of
- 4sU-tagged RNA at the MBT and the mid-gastrula stage. Numbers below RNAPII heat map
- represent counts of active (blue) and activated (red) genes at the indicated developmental
- stages. The horizontal dotted line separates RNAPII-engaged (RNAPII<sup>+</sup>) from non-engaged
- (RNAPII<sup>-</sup>) genes as detected until the late gastrula stage. The vertical dotted lines in the total
- RNA plots indicate the developmental time points of the MBT (white) and the late gastrula stage
- 625 (black), respectively.
- 626 (C) 2D space-filling (Hilbert) curves showing RNAPII recruitment to chromosomes (Chr) at the
- 128-cell stage and the MBT. A few zygotic genes are highlighted as being active (ON) or not
- 628 (OFF) based on their engagement with RNAPII.
- 629 (D) Alluvial diagram of spatio-temporal ZGA. Tissue-specificity inferred from regional transcript
- enrichments along the animal-vegetal or the dorso-ventral axes or both (Blitz et al., 2017).
- (E) ZGA-associated enrichment of biological processes.
- 632 (F) Length of maternal and/or zygotic genes. \*, p <1.9e-7 (Wilcoxon rank-sum test against
- maternal and post-MBT activated genes); r<sub>effect</sub> 0.06 ('MBT' vs 'Egg') 0.48 ('128' vs 'mG').
- 634 (G) Maternal/zygotic contribution to the transcriptome deduced from full-length RNAPII
- occupancy and maternally inherited RNA.
- Abbreviations: 32, 32-cell; 128, 128-cell; 1K, 1,024-cell; MBT, mid-blastula transition; mG, mid-
- gastrula; IG, late gastrula; 4sU, 4-thiouridine; Mdn, median; TPM, transcripts per million.
- See also Figure S1 and Tables S1 and S2.

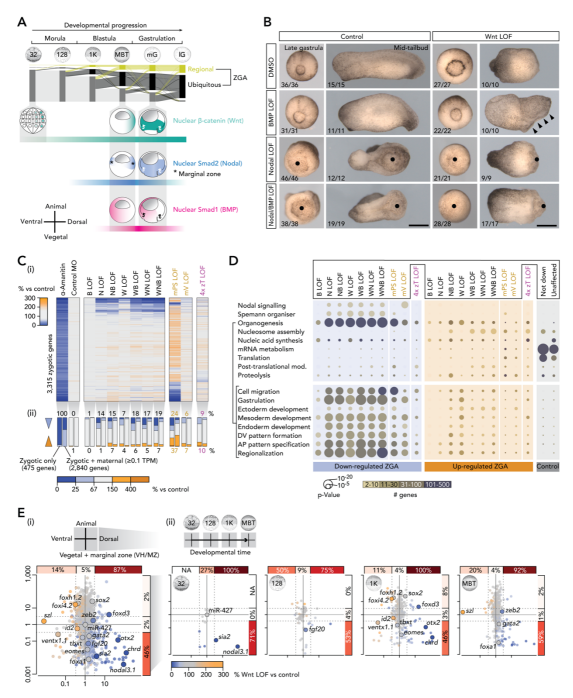


Figure 2. Spatio-temporal ZGA Regulated by Canonical Wnt, Nodal and BMP Signals

- (A) Spatio-temporal ZGA and nuclear localization of signal mediators  $\beta$ -catenin (canonical Wnt), Smad2 (Nodal) and Smad1 (BMP) (Faure et al., 2000; Larabell et al., 1997; Schohl and Fagotto, 2002) from the 32-cell to the late gastrula stage.
- (B) Morphological phenotypes of single and combined signal LOFs at the late gastrula and the mid-tailbud stage. Left ('control') pictures are taken from Gentsch et al. (2018b). Bullet points, failed blastopore formation. Arrowheads, excessive neural fold formation. Scale bar, 0.5 mm.
- (C) Heat map (i) and bar graph summary (ii) of ZGA mis-regulated in various LOF embryos: α-amanitin, positive control; control MO, negative control. Abbreviations: B, BMP; N, Nodal; W, canonical Wnt; mPS, maternal Pou5f3/Sox3; mV, maternal VegT; 4x zT, four zygotic T-box TFs (zygotic VegT, Eomes, Tbxt and Tbxt.2).

(D) Biological processes enriched with mis-regulated and control (not down-regulated or unaffected by the loss of maternal TFs or signaling) sets of zygotic genes under indicated LOFs.

(E) Summary (i) and temporal resolution (ii) of Wnt LOF effects on regional ZGA. Percentages only refer to the down-regulated genes among all zygotic genes with similar expression ratios along the animal-vegetal or the dorso-ventral axes.

See also Figure S2, Tables S1 and S3, and Movie S1.

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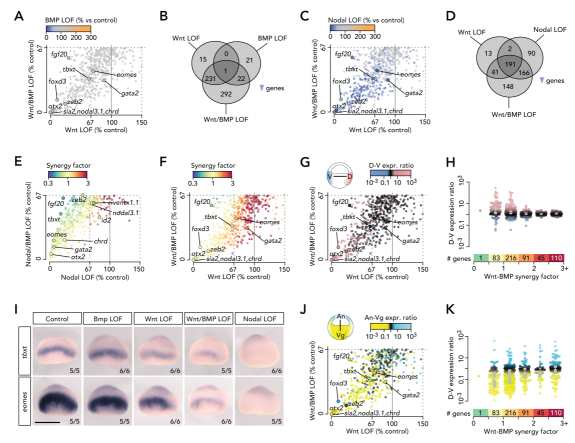


Figure 3. Uniform Gene Expression Across Dorso-Ventral Axis Achieved by Wnt/BMP Synergy.

(A,C,E,F,G,J) Scatter plots of relative (% to control) transcript levels between indicated LOFs with each dot (gene) color-coded according to a third attribute: (A,B) relative (% to control) transcript levels, (E,F) synergy factors between single inductive signals, and (G,J) regional expression ratios between opposite ends of the indicated axis.

- (B,D) Venn diagram of down-regulated genes by indicated LOFs.
- 665 (H,K) Box and beeswarm plots of regional expression (as measured along the indicated axes)
  666 depending on increased Wnt-BMP synergy.
- 667 (I) WMISH of *tbxt* and *eomes* transcripts under various LOFs. Control and Nodal LOF pictures are from Gentsch et al. (2018b).
- See also Figure S3 and Table S3.

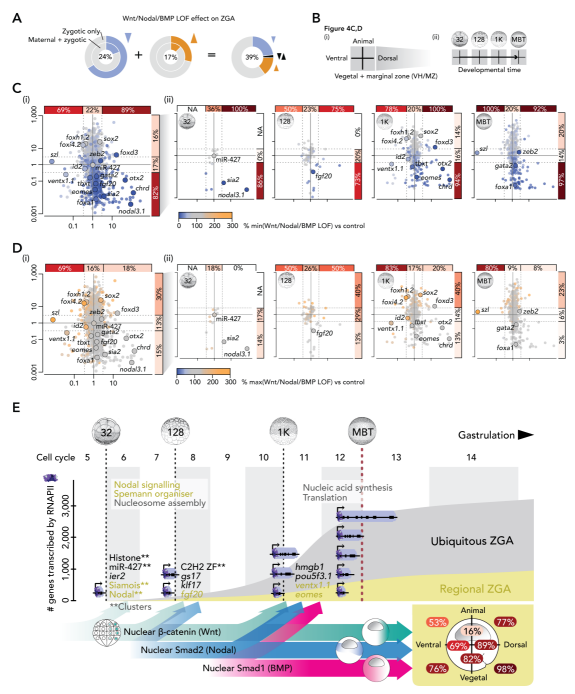


Figure 4. Canonical Wnt, Nodal and BMP Signals Induce the Majority of Regional ZGA.

- (A) Total percentage of active (zygotic only and maternal-zygotic) genes mis-regulated by Wnt, Nodal and/or BMP LOF.
- (B) Graphical explanations of figure panels (C) and (D).

- (C,D) Summary (i) and temporal resolution (ii) of minimal (C) and maximal (D) transcript levels of active genes (separated by regional expression along the primary body axes) detected among Wnt, Nodal and/or BMP LOFs. Percentages only refer to the down-regulated (C) or upregulated (D) genes among all zygotic genes with the same range of expression ratios along the animal-vegetal or dorso-ventral axes.
- (E) Exponential activation of gradually longer genes before the MBT (bulk ZGA) when cell divisions occur at rapid and nearly constant intervals (~20 min at 28°C). Sequential induction

of the canonical Wnt, Nodal and BMP pathway is critical to high percentages of regional ZGA (as measured along the two primary body axes within the indicated halves and quadrants of an early gastrula embryo): e.g. ~89% or ~98% of gene expression enriched in the dorsal half or vegetal-dorsal quadrant, respectively.

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## SUPPLEMENTAL INFORMATION

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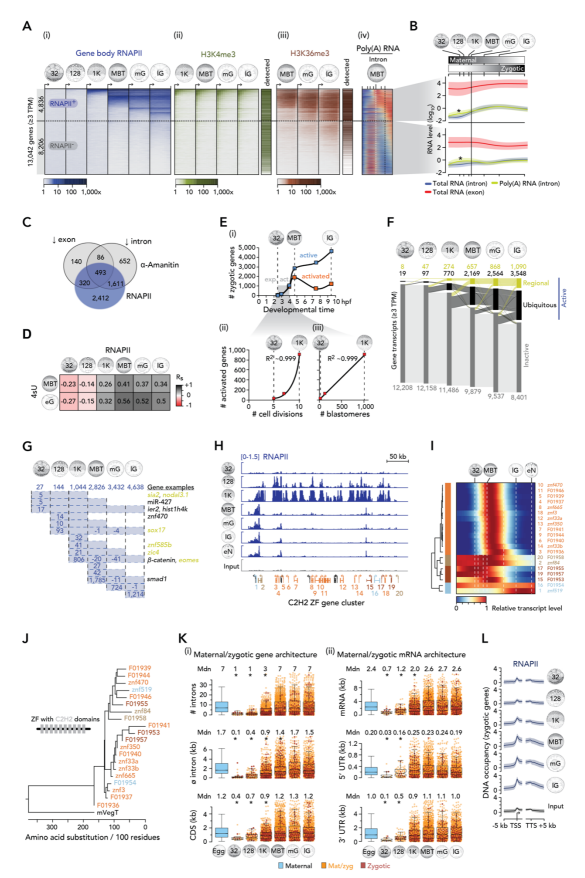


Figure S1. Dynamics and Architecture of ZGA in X. tropicalis, Related to Figure 1.

- (A) Progression of ZGA from the 32-cell to the late gastrula stage based on (i) whole gene body
- 690 (full-length) occupancy of RNAPII (i.e., RNAPII was enriched across entire gene bodies; see
- Transparent Methods). Co-aligned: Active histone marks H3K4me1 (ii) and H3K36me3 (iii)
- 692 (Hontelez et al., 2015) and intronic signal from the high time-resolution profile of poly(A) RNA
- (iv) (Owens et al., 2016). The horizontal dotted line separates RNAPII-engaged (RNAPII<sup>+</sup>) from
- non-engaged (RNAPII) genes as detected until the late gastrula stage. The vertical dotted lines
- in the poly(A) RNA plot indicate the developmental time points of the MBT (white) and the late
- gastrula stage (black), respectively.
- 697 (B) Transcript feature levels (mean +/- SD) during the maternal-to-zygotic transition. Asterisk,
- 698 polyadenylation immediately after fertilization (Collart et al., 2014) transiently increased the
- intronic signal obtained from the poly(A) RNA samples.
- (C) Venn diagram of zygotic genes detected by full-length RNAPII occupancy or reduced exonic
- 701 or intronic transcript counts upon blocking RNAPII-mediated transcription with  $\alpha$ -amanitin.
- 702 (D) Pairwise Spearman's correlations (R<sub>s</sub>) of enrichment values resulting from RNAPII profiling
- and 4sU tagging to detect zygotic genes at indicated developmental stages.
- (E) Plots of the number of active and newly activated genes (i) or newly activated genes versus
- the developmental time (i), the number of completed cell divisions (ii) or formed blastomeres
- 706 (iii).
- 707 (F) Alluvial diagram of spatio-temporal ZGA including maternally inherited RNA transcripts of
- 708 genes not activated by the late gastrula stage. Tissue-specificity inferred from regional
- 709 transcript enrichments along the animal-vegetal or the dorso-ventral or both axes (Blitz et al.,
- 710 2017).
- 711 (G) Numbers of genes with full-length RNAPII occupancy at indicated developmental stages.
- 712 Examples of ubiquitously (black) and tissue-specifically (orange) expressed genes are listed to
- 713 the right.
- 714 (H) RNAPII dynamics at the Cys2-His2 [C2H2] zinc finger (ZF) cluster from the 32-cell to the
- 715 early neurula stage.
- 716 (I) Expression dynamics of C2H2 ZF genes normalized to maximal transcript levels recorded
- 517 between fertilization and 23.5 hpf (Owens et al., 2016).
- 718 (J) Phylogenetic tree of the C2H2 ZF genes shown in (H,I). Maternal VegT (mVegT), outgroup
- 719 TF of this phylogenetic tree.
- 720 (K) Box and beeswarm plots showing various metrics of the zygotic/maternal genes (i) and
- 721 mRNA (ii) during ZGA. Asterisks, significant Wilcoxon rank-sum tests against maternal and
- 722 post-MBT activated genes and corresponding effect sizes ( $r_{effect}$ ): # introns, p <2.1e-13,  $r_{effect}$
- 723 0.07-0.43; Ø intron (kb), p <7.9e-7,  $r_{effect}$  0.07-0.42; CDS (kb), p <3e-7,  $r_{effect}$  0.05-0.27; mRNA
- 724 (kb), p <7e-11,  $r_{effect}$  0.06-0.32; 5' UTR (kb), p <0.015,  $r_{effect}$  0.03-0.15; and 3' UTR (kb), p <1.4e-
- 725 6, r<sub>effect</sub> 0.04-0.21.
- 726 (L) Meta-profiles (mean +/- SD) of RNAPII (separated by developmental stage) and input
- 727 (negative control) densities at zygotic genes.

- Abbreviations: 32, 32-cell; 128, 128-cell; 1K, 1,024-cell; MBT, mid-blastula transition; mG, mid-
- gastrula; IG, late gastrula; eN, early neurula; 4sU, 4-thiouridine; Mdn, median; TPM, transcripts
- 730 per million.

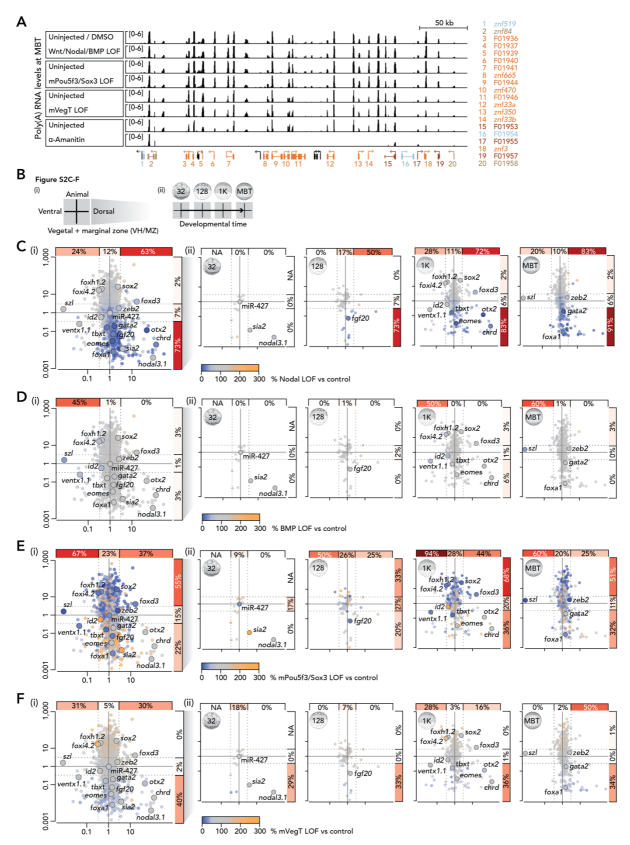


Figure S2. Effect of Canonical Wnt, Nodal and BMP Signals on ZGA, Related to Figure 2.

- (A) Poly(A) RNA profiles of the C2H2 ZF cluster (Figure S1H) for indicated control and LOFs at the MBT.
- 735 (B) Graphical explanations of figure panels (C-F).

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(C-F) Summary (i) and temporal resolution (ii) of gene mis-regulations upon the LOF of Nodal (C) or BMP (D) signaling or maternal Pou5f3/Sox3 (mPou5f3/Sox3) (E) or VegT (mVegT) (F). Percentages only refer to the down-regulated genes (by ≥1/3 compared to control expression level) among all zygotic genes with the same range of expression ratios along the animal-vegetal or the dorso-ventral axes.

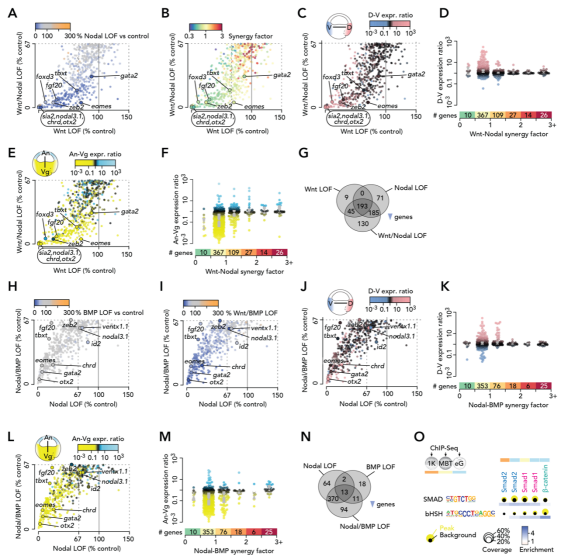


Figure S3. Relationship between Canonical Wnt, Nodal and BMP to Control Regional ZGA, Related to Figure 3.

(A-C,E,H-J,L) Scatter plots of relative (% to control) transcript levels between indicated LOFs with each dot (gene) color-coded according to a third attribute: (A,H,I) relative (% to control) transcript levels, (B) synergy factors between single inductive signals, and (C,E,J,L) regional expression ratios between opposite ends of the indicated axis.

- (G,N) Venn diagram of down-regulated genes by indicated LOFs.
- 749 (D,F,K,M) Box and beeswarm plots of regional expression (as measured along the indicated axes) depending on increased Wnt-Nodal (D,F) or Nodal-BMP (K,M) synergy.
  - (O) Coverage and enrichment of Smad and β-catenin-associated DNA motifs (SMAD and

bHSH motifs) at endogenous binding sites of β-catenin, Smad1, Smad2 (Gentsch et al., 2018b) at indicated developmental stages (color-coded).

## **SUPPLEMENTAL TABLES**

# Table S1. Summary of Deep sequencing and Read Alignments, Related to Figure 1 and 2.

The meta-data of ChIP-Seq, 4sU-Seq and RNA-Seq runs includes the developmental stage, condition, read type, and total and genome-aligned read numbers. \*, all rows represent manually collected biological sample: a, b and c mark biological replicates. †, library was sequenced twice. Reads were pooled from both sequencing runs. ¶, reported numbers are reads that are non-redundantly aligned to the genome assembly 7.1 with a mapping quality (MAPQ) of 10 (DNA) and 255 (RNA). NF, staging according to Nieuwkoop and Faber (1994). PE/SExx, paired-end or single-end sequencing plus read length.

Sample*	Library	Stage	NF	Condition	PE/SExx	Total reads	Aligned reads¶
1†	RNAPII ChIP	32-cell	6	None	SE50	45,073,468	25,451,597
2†	RNAPII ChIP	128-cell	7	None	SE50/100	65,757,131	41,352,403
3†	RNAPII ChIP	1,024-cell	8	None	SE50	60,445,294	37,661,064
4	RNAPII ChIP	MBT	8.5	None	SE50	29,196,397	18,402,411
5	RNAPII ChIP	mid-gastrula	11	None	SE50	29,268,648	20,085,357
6†	RNAPII ChIP	late gastrula	12+	None	SE50	95,959,570	67,591,256
7	RNAPII ChIP	early neurula	13+	None	SE50	47,039,656	32,889,013
8	4sU-tagged total RNA	Mid-blastula	8+	4sU (4-thiouridine)	PE50	22,202,216	2,738,397
9	4sU-tagged total RNA	Early gastrula	10+	4sU (4-thiouridine)	PE50	20,388,030	5,212,782
10	Total RNA	Mid-blastula	8+	4sU (4-thiouridine), input	PE50	15,087,016	11,451,545
11	Total RNA	Early gastrula	10+	4sU (4-thiouridine), input	PE50	14,902,023	9,771,572
12a	Poly(A)+ RNA	Late blastula	9+	Wnt LOF (β-catenin MO)	PE76	39,439,343	35,540,116
12b	Poly(A)+ RNA	Late blastula	9+	Wnt LOF (β-catenin MO)	PE76	53,523,555	48,364,696
12c	Poly(A)+ RNA	Late blastula	9+	Wnt LOF (β-catenin MO)	PE76	50,568,196	45,534,243
13a	Poly(A)+ RNA	Late blastula	9+	Wnt/Nodal LOF (β-catenin MO, SB431542)	PE76	43,980,513	39,732,439
13b	Poly(A)+ RNA	Late blastula	9+	Wnt/Nodal LOF (β-catenin MO, SB431542)	PE76	44,582,821	40,373,045
13c	Poly(A)+ RNA	Late blastula	9+	Wnt/Nodal LOF (β-catenin MO, SB431542)	PE76	37,243,519	33,648,196
14a	Poly(A)+ RNA	Late blastula	9+	Wnt/BMP LOF (β-catenin MO, LDN193189)	PE76	48,440,801	44,080,313
14b	Poly(A)+ RNA	Late blastula	9+	Wnt/BMP LOF (β-catenin MO, LDN193189)	PE76	40,989,331	37,188,245
14c	Poly(A)+ RNA	Late blastula	9+	Wnt/BMP LOF (β-catenin MO, LDN193189)	PE76	42,494,161	38,527,873
15a	Poly(A)+ RNA	Late blastula	9+	Wnt/Nodal/BMP LOF (β-catenin MO, SB431542, LDN193189)	PE76	52,761,134	47,815,313
15b	Poly(A)+ RNA	Late blastula	9+	Wnt/Nodal/BMP LOF (β-catenin MO, SB431542, LDN193189)	PE76	55,128,794	50,165,817
15c	Poly(A)+ RNA	Late blastula	9+	Wnt/Nodal/BMP LOF (β-catenin MO, SB431542, LDN193189)	PE76	57,414,981	52,101,210
16a	Poly(A)+ RNA	Late blastula	9+	Control (uninjected, DMSO)	PE76	42,248,469	37,989,215
16b	Poly(A)+ RNA	Late blastula	9+	Control (uninjected, DMSO)	PE76	52,066,825	47,105,847
16c	Poly(A)+ RNA	Late blastula	9+	Control (uninjected, DMSO)	PE76	42,875,722	38,893,706

Total						2,948,234,870	2,524,336,789
25d	Poly(A)+ RNA	Late gastrula	12+	Control (standard control MO)	PE76	39,860,708	35,323,477
25c	Poly(A)+ RNA	Late gastrula		Control (standard control MO)	PE76	48,184,154	42,775,598
25b	Poly(A)+ RNA	Late gastrula		Control (standard control MO)	PE76	36,043,121	31,918,744
25a	Poly(A)+ RNA	Late gastrula		Control (standard control MO)	PE76	53,566,350	47,558,871
24d	Poly(A)+ RNA	Mid-gastrula	11+	Control (standard control MO)	PE76	69,504,876	61,831,845
24c	Poly(A)+ RNA	Mid-gastrula	11+	Control (standard control MO)	PE76	56,633,663	50,510,544
24b	Poly(A)+ RNA	Mid-gastrula			PE76	54,847,378	48,981,425
24a	Poly(A)+ RNA	Mid-gastrula	11+	Control (standard control MO)	PE76	50,011,338	44,604,698
23d	Poly(A)+ RNA	Early gastrula		Control (standard control MO)	PE76	67,143,066	60,397,172
23c	Poly(A)+ RNA	Early gastrula		Control (standard control MO)	PE76	78,605,062	70,838,519
23b	Poly(A)+ RNA	Early gastrula		Control (standard control MO)	PE76	49,193,836	44,649,709
23a	Poly(A)+ RNA	Early gastrula	10+	(zVegT, Eomes, Tbxt, Tbxt.2 MO) Control (standard control MO)	PE76	57,881,388	52,311,548
22d	Poly(A)+ RNA	Late gastrula	12+	(zVegT, Eomes, Tbxt, Tbxt.2 MO) 4x zT (4 zygotic T-box TFs) LOF	PE76	56,546,178	49,962,419
22c	Poly(A)+ RNA	Late gastrula	12+	(zVegT, Eomes, Tbxt, Tbxt.2 MO) 4x zT (4 zygotic T-box TFs) LOF	PE76	54,924,184	48,697,757
22b	Poly(A)+ RNA	Late gastrula	12+	4x zT (4 zygotic T-box TFs) LOF	PE76	76,564,052	67,801,662
22a	Poly(A)+ RNA	Late gastrula	12+	4x zT (4 zygotic T-box TFs) LOF (zVegT, Eomes, Tbxt, Tbxt.2 MO)	PE76	59,905,689	53,050,487
21d	Poly(A)+ RNA	Mid-gastrula	11+	4x zT (4 zygotic T-box TFs) LOF (zVegT, Eomes, Tbxt, Tbxt.2 MO)	PE76	57,421,959	51,207,446
21c	Poly(A)+ RNA	Mid-gastrula	11+	4x zT (4 zygotic T-box TFs) LOF (zVegT, Eomes, Tbxt, Tbxt.2 MO)	PE76	57,420,101	51,441,091
21b	Poly(A)+ RNA	Mid-gastrula	11+	4x zT (4 zygotic T-box TFs) LOF (zVegT, Eomes, Tbxt, Tbxt.2 MO)	PE76	49,856,218	44,780,174
21a	Poly(A)+ RNA	Mid-gastrula	11+	4x zT (4 zygotic T-box TFs) LOF (zVegT, Eomes, Tbxt, Tbxt.2 MO)	PE76	60,071,724	53,786,168
20d	Poly(A)+ RNA	Early gastrula		4x zT (4 zygotic T-box TFs) LOF (zVegT, Eomes, Tbxt, Tbxt.2 MO)	PE76	53,693,870	48,621,416
				(zVegT, Eomes, Tbxt, Tbxt.2 MO)			, ,
20c	Poly(A)+ RNA	Early gastrula		(zVegT, Eomes, Tbxt, Tbxt.2 MO) 4x zT (4 zygotic T-box TFs) LOF	PE76	66,378,689	60,011,884
20b	Poly(A)+ RNA	Early gastrula		(zVegT, Eomes, Tbxt, Tbxt.2 MO) 4x zT (4 zygotic T-box TFs) LOF	PE76	62,453,340	56,799,027
20a	Poly(A)+ RNA	Early gastrula		LDN193189) 4x zT (4 zygotic T-box TFs) LOF	PE76	70,597,621	63,789,737
19c	Poly(A)+ RNA	Late blastula	9+	LDN193189) Nodal/BMP LOF (SB431542,	PE76	49,397,754	44,710,470
19b	Poly(A)+ RNA	Late blastula	9+	LDN193189) Nodal/BMP LOF (SB431542,	PE76	48,875,533	44,167,262
19a	Poly(A)+ RNA	Late blastula	9+	Nodal/BMP LOF (SB431542,	PE76	49,865,847	45,127,683
18c	Poly(A)+ RNA	Late blastula	9+	Bmp LOF (LDN193189)	PE76	40,645,238	36,899,368
18b	Poly(A)+ RNA	Late blastula	9+	Bmp LOF (LDN193189)	PE76	35,247,462	31,937,946
18a	Poly(A)+ RNA	Late blastula	9+	Bmp LOF (LDN193189)	PE76	49,171,585	44,430,698
17c	Poly(A)+ RNA	Late blastula	9+	Nodal LOF (SB431542)	PE76	47,497,608	43,105,868
17b	Poly(A)+ RNA	Late blastula	9+	Nodal LOF (SB431542)	PE76	43,201,587	39,270,622
17a	Poly(A)+ RNA	Late blastula	9+	Nodal LOF (SB431542)	PE76	47,946,077	43,367,783

# 764 Table S2. Temporal Progression of ZGA, Related to Figure 1.

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Activated genes are listed according to the earliest developmental stage of full-length RNAPII occupancy. The lists also include the overall enrichment of RNAPII across the gene body, the maternal contribution (TPM) (Owens et al., 2016) and the enrichment of de novo synthesized transcripts determined by 4sU tagging at the MBT and the early gastrula stage.

# Table S3. Differential ZGA Analysis, Related to Figure 2, 3 and 4.

- 770 Differential expression analysis of genes showing ≥50% (FDR ≤10%) reduced transcript levels
- 771 in α-amanitin-injected embryos. Normalized transcript levels (inferred from exon or intron
- counts) are scaled (percentage, %) to the expression level in control embryos. The list also
- includes FDRs, expression ratios across the animal-vegetal or dorso-ventral axis (Blitz et al.,
- 774 2017), the earliest developmental stage of full-length RNAPII occupancy, and average
- expression levels between 0 and 1 hpf (maternal) and 5 and 9 hpf (from the MBT to the mid-
- 776 gastrula stage).
- 777 Movie S1. Quadruple LOF of Zygotic T-box TFs, Related to Figure 2.
- 778 Simultaneous filming of the vegetal (top row) and animal (bottom row) hemisphere of 4x zT
- The LOF (labelled as T-box KD in the movie) (left) and control (right) embryos from early gastrula
- 780 to mid-tailbud stage.

- TRANSPARENT METHODS
- 782 CONTACT FOR REAGENTS AND RESOURCE SHARING
- Further information and requests for resources and reagents should be directed to and will be
- fulfilled by the Lead Contact, James C. Smith (jim.smith@crick.ac.uk).
- 785 EXPERIMENTAL MODEL AND SUBJECT DETAILS
- 786 Xenopus tropicalis Manipulation
- Standard procedures were used for ovulation, fertilization, and manipulation and incubation of
- 788 embryos (Khokha et al., 2002; Sive et al., 2000). Briefly, frogs were obtained from Nasco
- 789 (Wisconsin, USA). Ovulation was induced by injecting serum gonadotropin (Intervet) and
- 790 chorionic gonadotropin (Intervet) into the dorsal lymph sac of mature female frogs. Eggs were
- 791 fertilized *in vitro* with sperm solution consisting of 90% Leibovitz's L-15 medium (Thermo Fisher
- 792 Scientific, Cat#11415064) and 10% fetal bovine serum (Thermo Fisher Scientific,
- 793 Cat#10500056). After 10 min, fertilized eggs were de-jellied with 2.2% (w/v) L-cysteine (Merck,
- 794 Cat#168149) equilibrated to pH 8.0. Embryos were cultured in 5% Marc's Modified Ringer's
- solution (MMR) (5 mM NaCl, 0.1 mM KCl, 0.1 mM CaCl<sub>2</sub>, 0.05 mM MgSO<sub>4</sub> and 0.25 mM HEPES
- 796 pH7.5) at 21°C-28°C. Embryos were staged according to Nieuwkoop and Faber (1994). All
- 797 Xenopus work fully complied with the UK Animals (Scientific Procedures) Act 1986 as
- 798 implemented by the Francis Crick Institute.
- 799 Chromatin immunoprecipitation (ChIP)
- 800 ChIP was carried out as detailed previously (Gentsch and Smith, 2017). Briefly, de-jellied X.
- tropicalis embryos were fixed at room temperature with 1% formaldehyde (Merck, Cat#F8775)
- in 1% MMR for 25 min. The fixation time was extended to 45 min for pre-gastrula stages. The
- following number of embryos were used for ChIP-Seq: 1,400 at the 32-cell stage, 1,000 at the

128-cell stage, 700 at the 1,024-cell stage, 450 at the MBT and 350 for the post-MBT stages. Fixation was terminated by rinsing embryos three times with ice-cold 1% MMR. Fixed embryos were homogenized in CEWB1 (150 mM NaCl, 1 mM EDTA, 1% (v/v) Igepal CA-630 [Merck, Cat#I3021], 0.25% (w/v) sodium deoxycholate [Merck, Cat#SRE0046], 0.1% (w/v) sodium dodecyl sulfate [Merck, Cat#71729] and 10 mM Tris-HCl pH 8.0) supplemented with 0.5 mM **DL-Dithiothreitol** (Fluorochem, Cat#M02712) and protease inhibitors (Roche, Cat#11873580001). The homogenate was left on ice for 5 min and then centrifuged at 1,000 g (4°C) for 5 min. Homogenization and centrifugation was repeated once before resuspending the pellet in 1-3 ml CEWB1. Chromatin was solubilized and fragmented by microtip-mediated ultra-sonication (Misonix 3000 sonicator with a tapered 1/16-inch microtip). The solution of fragmented chromatin was cleared by centrifuging at 16,000 g (4°C) for 5 min. About 1% of the cleared chromatin extract was set aside for the input sample (negative control). The remaining chromatin was incubated overnight at 4°C on a vertical rotor (10 rpm) with 20 µl of the mouse monoclonal anti-RNAPII (8WG16) (Covance, Cat#MMS-126R; RRID: AB 10013665) antibody. After adding 100 µl of washed protein G magnetic beads (Thermo Fisher Scientific, Cat#10003D) the solution was incubated for another 4 h at 4°C on a vertical rotor (10 rpm). The beads were washed eight times in CEWB1 and once in TEN (10 mM Tris-HCl pH 8.0, 150 mM NaCl and 1 mM EDTA) at 4°C. ChIP was eluted off the beads twice with 100 µl SDS elution buffer (50 mM Tris-HCl pH 8.0, 1 mM EDTA and 1% (w/v) sodium dodecyl sulfate) at 65°C. ChIP eluates were pooled before reversing DNA-protein cross-links. Input (filled up to 200 µl with SDS elution buffer) and ChIP samples were supplemented with 10 µl 5 M NaCl and incubated at 65°C for 6-16 h. Samples were treated with proteinase K (Thermo Fisher Scientific, Cat#AM2548) and RNase A (Thermo Fisher Scientific, Cat#12091021) to remove any proteins and RNA from the co-immunoprecipitated DNA fragments. The DNA was purified with phenol:chloroform:isoamyl alcohol (25:24:1, pH 7.9) (Thermo Fisher Scientific, Cat#AM9730) using 2.0-ml Phase Lock Gel Heavy microcentrifuge tubes (VWR, Cat#733-2478) for phase separation and precipitated with 1/70 volume of 5 M NaCl, 2 volumes of absolute ethanol and 15 μg GlycoBlue (Thermo Fisher Scientific, Cat#AM9516). After centrifugation, the DNA pellet was air-dried and dissolved in 11 µl elution buffer (10 mM Tris-HCl pH 8.5). The DNA concentration was determined on a Qubit fluorometer using high-sensitivity reagents for detecting double-stranded DNA (10 pg/µl to 100 ng/µl) (Thermo Fisher Scientific, Cat#Q33231).

## **ChIP-Seq Library Preparation**

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Using the KAPA Hyper Prep Kit (Roche, Cat#KK8504), 2.5-5 ng ChIP DNA or 5 ng input DNA were converted into indexed paired-end libraries as previously described (Gentsch and Smith, 2017). Briefly, DNA fragments were end-repaired and A-tailed for 30 min at 20°C followed by 30 min at 65°C before cooling to 4°C. 7.5 pmol TruSeq (single index) Y-adapters (IDT) were ligated to the DNA fragment ends for 20 min at 20°C. The DNA ligation product was extracted with 0.8x SPRI (solid phase reversible immobilisation) beads (Beckman Coulter, Cat#A63882) and amplified in five PCR cycles (15 sec at 98°C, 30 sec at 60°C and 30 sec at 72°C) using the KAPA high-fidelity

polymerase master mix (Roche, Cat#KK2602) and 25 pmol Illumina P5 (forward) and P7 (reverse) primers (IDT). After cleaning up the PCR reaction with 1x SPRI beads, the DNA library was size-separated by electrophoresis using E-gel EX agarose gels (Thermo Fisher Scientific, Cat#G401002). A gel slice containing DNA ranging from 250 to 450 bp in size was dissolved shaking in 350 μl QG buffer (Qiagen) using a thermomixer (1,000 rpm) at room temperature. The DNA was purified with MinElute columns (Qiagen, Cat#28604) and eluted off these columns twice using 11 μl elution buffer (10 mM Tris-HCl pH 8.5). The library was re-amplified using another 6-8 PCR cycles yielding 100-200 ng DNA without adapter dimer contamination. The DNA library was cleaned up with 1x SPRI beads.

## Illumina Sequencing

All sequencing libraries were quality controlled: The DNA yield and fragment size distribution were determined by fluorometry and chip-based capillary electrophoresis, respectively. ChIP-Seq and RNA-Seq libraries were sequenced on the Illumina HiSeq 2500 and 4000, respectively, by the Advanced Sequencing Facility of the Francis Crick Institute. Sequencing samples and read alignment results are summarized in Table S1.

# Post-Sequencing Analysis of ChIP-Seq

Single reads of maximal 50 bases were processed using trim\_galore v0.4.2 (Babraham Institute, UK) to trim off low-quality bases (default Phred score of 20, i.e. error probability was 0.01) and adapter contamination from the 3' end. Processed reads were aligned to the *X. tropicalis* genome assembly v7.1 and v9.1 (for Hilbert curves) running Bowtie2 v2.2.9 (Langmead and Salzberg, 2012) with default settings (Table S1). Alignments were converted to the HOMER's tag density format (Heinz et al., 2010) with redundant reads being removed (makeTagDirectory -single -tbp 1 -unique -mapq 10 -fragLength 175 -totalReads all). Only uniquely aligned reads (i.e. MAPQ ≥10) were processed. We pooled all input alignments from various developmental stages (Gentsch et al., 2018b). This created a comprehensive mappability profile that covered ~400 million unique base pair positions. For Hilbert curves, tag densities were generated across the genome v9.1 using sliding (200-bp increments) 400-bp window. Background signals (<0.3 reads per 1 million mapped reads) were removed. Blacklisted (Gentsch et al., 2018b) regions (except for MIR-427) were excluded using intersectBed (-v -f 0.5) from BEDtools v2.25.0 (Quinlan and Hall, 2010).

# Detecting Zygotic and Maternal Genes Using RNAPII Profiling and High Time-Resolution Transcriptomics

Normalized RNAPII and input tag densities were calculated across the gene body in 10 bins of equal size. Gene annotations v7.1 were altered based on a few known zygotic isoforms and some corrections obtained from assembling total and poly(A) RNA (Owens et al., 2016) from stage 6 to stage 12.5 *de novo* (Pertea et al., 2016). A few genes had previously been annotated as gene clusters due to assembly uncertainties. We reduced the annotation of polycistronic MIR-427 to the minus arm (scaffold\_3b:3516900-3523400) and only monitored *nodal3.5* and *nodal5.3* within their

respective gene clusters. Gene bodies with <40% mappability were removed. Here, the threshold of mappability per bin was set at 10% of the input read density averaged across all gene bodies in use. Subsequently, enrichment values were only obtained for all mappable bins by dividing read densities of RNAPII and input. Further, we restricted the analysis to genes for which ≥3 transcripts per million (TPM) could be detected on average over three consecutive time points (i.e. over the developmental time of 1 h) of a high-resolution profile of total RNA (Owens et al., 2016) from fertilization to after gastrulation (stage 13). Genes were considered active when RNAPII enrichments along their full length (see thresholds below) and corresponding transcripts (≥0.1 TPM) were simultaneously detected. Transcript levels were calculated over three consecutive time points +/- 1 h from the developmental stage of RNAPII profiling. RNAPII enrichment covered ≥80% of the mappable gene body and reached at least one of the following thresholds: (1) 2.6-fold, (2) 1.8-fold and 1.4-fold at the next or previous stage, (3) 1.4-fold and 1.8-fold at the next or previous stage, or (4) 1.4-fold over three consecutive stages. The heatmap (Figure 1B and S1A) was sorted by the developmental stage (1st) and the overall fold (2nd) of RNAPII enrichment. Zygotic and maternal contributions to transcriptome (Figure 1G) were based on RNAPII enrichment (see above) and mean transcript levels (≥0.1 TPM) detected between 0 and 1 hpf, respectively.

# **Peak Calling and Motif Enrichment Analysis**

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Peak calling and motif enrichment analysis were carried out as previously reported (Gentsch et al., 2018b). Briefly, HOMER v4.8.3 (Heinz et al., 2010) was used to identify the binding sites of Smad1 (Gentsch et al., 2018b), Smad2 (Chiu et al., 2014; Gentsch et al., 2018b; Yoon et al., 2011) and β-catenin (Gentsch et al., 2018b; Nakamura et al., 2016) by virtue of ChIP-enriched read alignments (hereafter called peaks): findpeaks -style factor -minDist 175 -fragLength 175 inputFragLength 175 -fdr 0.001 -gsize 1.435e9 -F 3 -L 1 -C 0.97. This means that both ChIP and input alignments were extended 3' to 175 bp for the detection of significant (FDR ≤0.1%) peaks being separated by ≥175 bp. The effective size of the X. tropicalis genome assembly v7.1 was set to 1.435 billion bp, an estimate obtained from the mappability profile (Gentsch et al., 2018b). These peaks showed equal or higher tag density than the surrounding 10 kb, ≥3-fold more tags than the input and ≥0.97 unique tag positions relative to the expected number of tags. To further eliminate any false positive peaks, we removed any peaks with <0.5 CPM and those falling into blacklisted regions showing equivocal mappability due to genome assembly errors, gaps or simple/tandem repeats. Regions of equivocal mappability were identified by a two-fold lower (poor) or three-fold higher (excessive) read coverage than the average detected in 400-bp windows sliding at 200-bp intervals through normalized ChIP input and DNase-digested naked genomic DNA (Gentsch et al., 2018b). All identified regions ≤800 bp apart were subsequently merged. Gap coordinates were obtained from the Francis Crick mirror site of the UCSC (http://genomes.crick.ac.uk). Simple repeats were masked with RepeatMasker v4.0.6 (Smit et al.) using the crossmatch search engine v1.090518 (Phil Green) and the following settings: RepeatMasker -species "xenopus silurana tropicalis" -s -xsmall. Tandem repeats were masked with Jim Kent's trfBig wrapper script of the Tandem Repeat Finder v4.09 (Benson, 1999) using the

924 following settings: weight for match, 2; weight for mismatch, 7; delta, 7; matching probability, 80;

indel probability, 10; minimal alignment score, 50; maximum period size, 2,000; and longest

tandem repeat array (-I), 2 [million bp]. The enrichment and occurrence of predetermined DNA

binding motifs was calculated using 100 bp centred across the top 2,000 peaks per chromatin

feature and developmental stage: findMotifsGenome.pl -size 100 -mknown -nomotif.

## **Injections and Treatments of Embryos**

- 931 Microinjections were performed using calibrated needles and embryos equilibrated in 4% (w/v)
- 932 Ficoll PM-400 (Merck, Cat#F4375) in 5% MMR. Microinjection needles were generated from
- 933 borosilicate glass capillaries (Harvard Apparatus, GC120-15) using the micropipette puller
- 934 Sutter p97. Maximally three nanolitres were injected into the animal hemisphere of de-iellied
- 935 zygotes using the microinjector Narishige IM-300. Embryos were transferred to fresh 5% MMR
- 936 (without Ficoll PM-400) once they reached about the mid-blastula stage.
- 937 For profiling the nascent transcriptome, embryos were injected with 75 ng 4-thiouridine-5'-
- 938 triphosphate (4sU) (TriLink BioTechnologies, Cat#N-1025), which is incorporated into newly
- 939 synthesized transcripts.

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- 940 Loss-of-functions (LOFs) were generated by treating embryos with small molecule inhibitors
- 941 and/or injecting them with morpholinos (MOs) or α-amanitin. MOs were designed and produced
- 942 by Gene Tools (Oregon, USA) to block splicing (MO<sub>splice</sub>) or translation (MO<sub>transl</sub>): maternal
- 943 Pou5f3/Sox3 (mPou5f3/Sox3) LOF, 5 ng Pou5f3.2 MO<sub>transl</sub> (Chiu et al., 2014; Gentsch et al.,
- 944 2018b; GCTGTTGGCTGTACATAGTGTC), 5 ng Pou5f3.3 MO<sub>transl</sub> (Chiu et al., 2014; Gentsch
- 945 et al., 2018b; TACATTGGGTGCAGGGACCCTCTCA) and 5 ng Sox3 MO<sub>transl</sub> (Gentsch et al.,
- 946 2018b; GTCTGTGTCCAACATGCTATACATC); maternal VegT (mVegT) LOF, 10 ng mVegT
- 947 MO<sub>transl</sub> (Gentsch et al., 2018b; Rana et al., 2006; TGTGTTCCTGACAGCAGTTTCTCAT);
- 948 canonical LOF, 5 ng β-catenin  $MO_{transl}$ (Heasman
- 949 TTTCAACAGTTTCCAAAGAACCAGG); LOF of four zygotic T-box TFs (4x zT LOF), 2.5 ng tbxt
- 950 (Xbra, t) MO<sub>solice</sub> (Gentsch et al., 2013; TGGAGAGACCCTGATCTTACCTTCC), 2.5 ng tbxt
- 951 (Xbra, t) MO<sub>transl</sub> (Gentsch et al., 2013; GGCTTCCAAGCGCACACTGGG), 2.5 ng tbxt.2
- 952 (Xbra3, t2) MO<sub>splice</sub> (Gentsch et al., 2013; GAAAGGTCCATATTCTCTTACCTTC), 2.5 ng tbxt.2
- 953 (Xbra3, t2) MO<sub>transl</sub> (Gentsch et al., 2013; AGCTGTGCCTGTGCTCATTGTATTG), 5 ng zVegT
- 954 MO<sub>transl</sub> (Fukuda et al., 2010; Gentsch et al., 2013; CATCCGGCAGAGAGTGCATGTTCCT) and
- 955 5 eomes MO<sub>splice</sub> (Fukuda et al., 2010; Gentsch 2013;
- 956 GAACATCCTCCTGCAAAGCAAAGAC); control MO, 5-20 ng standard control MO
- 957 (CCTCTTACCTCAGTTACAATTTATA) according to the dose used for the β-catenin, mVegT
- 958 and 4x zT LOF experiment; and 30 pg α-amanitin (BioChemica, Cat#A14850001). To block
- 959 Nodal (Nodal LOF) and BMP (BMP LOF) signaling, embryos were treated with 100 µM
- 960 SB431542 (Tocris, Cat#1614) and/or 10 µM LDN193189 (Selleckchem, Cat#S2618) from the
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- 962 antagonists were dissolved. Transcriptional effects of combinatorial signal LOF were

8-cell stage onwards. Control embryos were treated accordingly with DMSO, in which these

determined at late blastula stage (stage 9<sup>+</sup>), while those of all other maternal LOFs were determined over three consecutive time points: the MBT (stage 8+), the late blastula (stage 9<sup>+</sup>) and the early gastrula (stage 10<sup>+</sup>) stage. The 4x zT LOF was transcriptionally profiled at early, mid and late gastrula stage (stage 10<sup>+</sup>, 11<sup>+</sup> and 12<sup>+</sup>). The 4x zT LOF comparison has four biological replicates (n=4). All other comparisons entail three biological replicates (n=3).

#### **Extraction of Total RNA**

Embryos were homogenized in 800 μl TRIzol reagent (Thermo Fisher Scientific, Cat#15596018) by vortexing. The homogenate was either snap-frozen in liquid nitrogen and stored at -80°C or processed immediately. For phase separation, the homogenate together with 0.2x volume of chloroform was transferred to pre-spun 2.0-ml Phase Lock Gel Heavy microcentrifuge tubes (VWR), shaken vigorously for 15 sec, left on the bench for 2 min and spun at ~16,000 g (4°C) for 5 min. The upper phase was mixed well with one volume of 95-100% ethanol and spun through the columns of the RNA Clean & Concentrator 25 Kit (Zymo Research, Cat#R1017) at ~12,000 g for 30 sec. Next, the manufacturer's instructions were followed for the recovery of total RNA (>17 nt) with minor modifications. First, the flow-through of the first spin was re-applied to the column. Second, the RNA was treated in-column with 3 U Turbo DNase (Thermo Fisher Scientific, Cat#AM2238). Third, the RNA was eluted twice with 25 μl molecular-grade water. The concentration was determined on the NanoDrop 1000 spectrophotometer or by fluorometry before depleting ribosomal RNA from total RNA (Profiling the Nascent Transcriptome).

# **Tagging the Nascent Transcriptome**

Thirty 4sU-injected embryos were collected at the MBT and the early-to-mid gastrula stage. Total RNA was extracted as outlined above. The 4sU-tagging was performed according to Gay et al. (2014) with few minor modifications. The RNA Clean & Concentrator 5 Kit (Zymo Research, Cat#R1013) was used to purify RNA. Briefly, the Ribo-Zero Gold rRNA Removal Kit (Illumina, Cat#MRZG126) was used according to the manufacturer's instructions to deplete ribosomal RNA from ~10 µg total RNA. The RNA was purified and fragmented for 4 min at 95°C using the NEBNext Magnesium RNA Fragmentation Module (NEB, Cat#E6150). The RNA was purified again before conjugating HDPD-Biotin (Thermo Fisher Scientific, Cat#21341) to 4sU via disulfide bonds for 3 h in the dark. Purified RNA was mixed with Streptavidin beads (Thermo Fisher Scientific, Cat#65305) to pull down biotin-tagged RNA. The RNA was eluted off the beads by treating them twice with 100 μl pre-heated (80°C) 100 mM β-mercaptoethanol (Merck, Cat#M6250), which breaks the disulfide bond between Biotin and 4sU. Subsequently, the RNA was converted into a deep sequencing library by following the manual instructions (Rev. C, 8/2014) of the ScriptSeq v2 RNA-Seq Library Preparation Kit (Illumina, Cat#SSV21106) starting with 4.1.A. (Anneal the cDNA Synthesis Primer) and 4.1.B. (Synthesize cDNA), RNA and ending with part 3.C (Synthesize 3'-Tagged DNA) to 3.G. (Assess Library Quantity and Quality). cDNA was purified using 1.8x SPRI beads. Input and 4sU-enriched cDNA were PCR-amplified with 11 and 15 cycles, respectively. The 4sU RNA-Seg library was purified with 1x SPRI beads.

# Post-Sequencing Analysis of 4sU Tagging

Paired-end reads were aligned to the *X. tropicalis* transcriptome assembly v7.1 running Bowtie2 (Langmead and Salzberg, 2012) with the following constraints: -k 200 (maximal allowed number of alignments per fragment) -X 800 (maximum fragment length in bp) --rdg 6,5 (penalty for read gaps of length N, 6+N\*5) --rfg 6,5 (penalty for reference gaps of length N, 6+N\*5) --score-main L,-.6,-.4 (minimal alignment score as a linear function of the read length x, f(x) = -0.6 - 0.4\*x) --no-discordant (no paired-end read alignments breaching maximum fragment length X) --no-mixed (only concordant alignment of paired-end reads). Only read pairs that uniquely align to one gene were counted. Raw read counts were normalized with DESeq2 v1.22.1 (Love et al., 2014) and then scaled to the input.

# Poly(A) RNA-Seq Profiling

10-15 embryos were collected per stage and condition. Total RNA was extracted as outlined above. Libraries were made from ~1  $\mu$ g total RNA by following the low-sample protocol of the TruSeq RNA Library Prep Kit v2 (Illumina, Cat#RS-122-2001) with a few modifications. First, 1  $\mu$ l cDNA purified after second strand synthesis was quantified on a Qubit fluorometer using high-sensitivity reagents for detecting double-stranded DNA (10  $\mu$ g/ $\mu$ l to 100  $\mu$ g/ $\mu$ l). By this stage, the yield was ~10  $\mu$ g. Second, the number of PCR cycles was reduced to eight to avoid products of over-amplification such as chimera fragments.

## Poly(A) RNA-Seq Read Alignment

Paired-end reads were aligned to the *X. tropicalis* genome assembly v7.1 using STAR v2.5.3a (Dobin et al., 2013) with default settings. The alignment was guided by a revised version of the gene models v7.2 (Collart et al., 2014) to improve mapping accuracy across splice junctions. The alignments were sorted by read name using the sort function of Samtools v1.3.1 (Li et al., 2009). Exon and intron counts (-t 'exon;intron') were extracted from unstranded (-s 0) alignment files using VERSE v0.1.5 (Zhu et al., 2016) in featureCounts (default) mode (-z 0). Intron coordinates were adjusted to exclude any overlap with exon annotation. For visualization, genomic BAM files of biological replicates were merged using Samtools and converted to the bigWig format. These genome tracks were normalized to the wigsum of 1 billion excluding any reads with mapping quality <10 using the python script bam2wig.py from RSeQC v2.6.4 (Wang et al., 2012).

## **Differential Gene Expression Analysis**

Differential expression analysis was performed with both raw exon and intron counts excluding those belonging to ribosomal and mitochondrial RNA using the Bioconductor/R package DESeq2 v1.22.1 (Love et al., 2014). In an effort to find genes with consistent fold changes over time, p-values were generated according to a likelihood ratio test reflecting the probability of rejecting the reduced (~ developmental stage) over the full (~ developmental stage + condition) model. Resulting p-values were adjusted to obtain false discovery rates (FDR) according to the Benjamini-Hochburg procedure with thresholds on Cook's distances and independent filtering being switched off. Equally, combinatorial LOF profiling and regional expression datasets (Blitz

et al., 2017) without time series were subjected to likelihood ratio tests with reduced (~ 1) and full (~ condition) models for statistical analysis. Fold changes of intronic and exonic transcript levels were calculated for each developmental stage and condition using the mean of DESeq2normalized read counts from biological replicates. Both intronic and exonic datasets were filtered for ≥10 DESeq2-normalized read counts that were detected at least at one developmental stage in all uninjected or DMSO-treated samples. Gene-specific fold changes were removed at developmental stages that yielded <10 normalized read counts in corresponding control samples. Next, the means of intronic and exonic fold changes were calculated across developmental stages. The whole dataset was confined to 3,318 genes for which at least 50% reductions (FDR ≤10%) in exonic (default) or intronic counts could be detected in a-amanitin-injected embryos. Regional expression was based on exonic read counts by default unless the intronic fold changes were significantly (FDR ≤10%) larger than the exonic fold changes (Table S3). For the hierarchical clustering of relative gene expression (Figure 2C), increased transcript levels were masked and only data points from signal LOFs, mPou5f3/Sox3 LOF and 4x zT LOF embryos were used. Euclidean distance-derived clusters were linked according to Ward's criterion and sorted using the optimal leaf ordering (OLO) algorithm. The synergy factor (SF) between signals x and y (Figures 3 and S3) were calculated as follows:  $SF_{xy} = \Delta_{xy} / (\Delta_x + \Delta_y)$ .  $\Delta$  is the relative loss of gene expression caused by signal depletion. For these calculations, any gene upregulations were neutralised (i.e. set to 1).

## Analysis of Enriched Gene Ontology (GO) Terms

- 1062 Over-represented GO terms were found by applying hypergeometric tests of the
- Bioconductor/R package GOstats v2.42.0 (Falcon and Gentleman, 2007) on gene lists. The
- process was also supported by the Bioconductor/R packages GSEABase v1.44.0 (Morgan et
- al., 2017) and GO.db v3.4.1 (Carlson et al., 2007). The gene universe was associated with GO
- terms by means of BLAST2GO (Conesa et al., 2005) as previously outlined (Collart et al., 2014;
- 1067 Gentsch et al., 2015).

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## **Generation of Hybridization Probes**

- Plasmids X. laevis eomes pCRII-TOPO (Gentsch et al., 2013) and X. laevis tbxt (Xbra, t) pSP73
- 1070 (Smith et al., 1991) were linearized by restriction digestion (BamHI and Bg/II, respectively) and
- purified using the QIAquick PCR Purification Kit (Qiagen, Cat#28104). The hybridization probes
- 1072 were transcribed from ~1 µg linearized plasmid using 1x digoxigenin-11-UTP (Roche,
- 1073 Cat#11277065910), 40 U RiboLock RNase inhibitor (Thermo Fisher Scientific, Cat#E00381), 1x
- transcription buffer (Roche) and T7 RNA polymerase (Roche, Cat#10881767001) at 37°C for 2 h.
- 1075 The probe was treated with 2 U Turbo DNase (Thermo Fisher Scientific) to remove the DNA
- 1076 template and purified by LiCl precipitation. RNA was diluted to 10 ng/µl (10x stock) with
- 1077 hybridization buffer. The hybridization buffer (stored at -20°C) consists of 50% formamide (Fisher
- Scientific, Cat#10052370), 5x saline sodium citrate (SSC), 1x Denhardt's solution (Thermo Fisher
- 1079 Scientific, Cat#750018), 10 mM EDTA, 1 mg/ml torula RNA (Merck, Cat#R6625), 100 μg/ml

heparin (Merck, Cat#H4784), 0.1% (v/v) Tween-20 (Merck, Cat#P9416) and 0.1% (w/v) CHAPS (Merck, Cat#C3023).

# Whole-Mount In Situ Hybridization (WMISH)

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WMISH was conducted using digoxigenin-labeled RNA probes (Monsoro-Burg, 2007; Sive et al., 2000). Briefly, X. tropicalis embryos were fixed in MEMFA (100 mM MOPS pH 7.4, 2 mM EDTA, 1 mM MgSO₄ and 3.7% formaldehyde) at room temperature for 1 h. The embryos were then washed once in 1x PBS and two to three times in ethanol. Fixed and dehydrated embryos were kept at -20°C for ≥24 h to ensure proper dehydration before starting hybridization. Dehydrated embryos were washed once more in ethanol before rehydrating them in two steps to PBT (1x PBS and 0.1% (v/v) Tween-20). Embryos were treated with 5 µg/ml proteinase K (Thermo Fisher Scientific) in PBT for 6-8 min, washed briefly in PBT, fixed again in MEMFA for 20 min and washed three times in PBT. Embryos were transferred into baskets, which were kept in an 8x8 microcentrifuge tube holder sitting inside a 10x10 slot plastic box filled with PBT. Baskets were built by replacing the round bottom of 2-ml microcentrifuge tubes with a Sefar Nitex mesh. This container system was used to readily process several batches of embryos at once. These baskets were maximally loaded with 40 to 50 X. tropicalis embryos. The microcentrifuge tube holder was used to transfer all baskets at once and to submerge embryos into subsequent buffers of the WMISH protocol. Next, the embryos were incubated in 500 µl hybridization buffer (see recipe above) for 2 h in a hybridization oven set to 60°C. After this prehybridization step, the embryos were transferred into 500 µl digoxigenin-labeled probe (1 ng/µl) preheated to 60°C and further incubated overnight at 60°C. The pre-hybridization buffer was kept at 60°C. The next day embryos were transferred back into the pre-hybridization buffer and incubated at 60°C for 10 min. Subsequently, they were washed three times in 2x SSC/0.1% Tween-20 at 60°C for 10 min, twice in 0.2x SSC/0.1% Tween-20 at 60°C for 20 min and twice in 1x maleic acid buffer (MAB) at room temperature for 5 min. Next, the embryos were treated with blocking solution (2% Blocking Reagent [Merck, Cat#11096176001] in 1x MAB) at room temperature for 30 min, and incubated in antibody solution (10% lamb serum [Thermo Fisher Scientific, Cat#16070096], 2% Blocking Reagent [Merck], 1x MAB and 1:2,000 Fab fragments from polyclonal anti-digoxigenin antibodies conjugated to alkaline phosphatase [Roche, Cat#11093274910; RRID:AB 514497]) at room temperature for 4 h. The embryos were then washed four times in 1x MAB for 10 min before leaving them in 1x MAB overnight at 4°C.

On the final day of the WMISH protocol, the embryos were washed another three times in 1x MAB for 20 min and equilibrated to working conditions of alkaline phosphatase (AP) for a total of 10 min by submerging embryos twice into AP buffer (50 mM MgCl<sub>2</sub>, 100 mM NaCl, 100 mM Tris-HCl pH 9.5 and 1% (v/v) Tween-20). At this stage, the embryos were transferred to 5-ml glass vials for monitoring the progression of the AP-catalyzed colorimetric reaction. Any residual AP buffer was discarded before adding 700  $\mu$ l staining solution (AP buffer, 340  $\mu$ g/ml nitro-blue tetrazolium chloride [Roche, Cat#11383213001] and 175  $\mu$ g/ml 5-bromo-4-chloro-3'-

indolyphosphate [Roche, Cat#11383221001]). The colorimetric reaction was developed at room temperature in the dark. Once the staining was clear and intense enough, the color reaction was stopped by two washes in 1x MAB. To stabilize and preserve morphological features, the embryos were fixed with Bouin's fixative without picric acid (9% formaldehyde and 5% glacial acetic acid [Fisher Scientific, Cat#10171460]) at room temperature for 30 min. Next, the embryos were washed twice in 70% ethanol/PBT to remove the fixative and residual chromogens. After rehydration to PBT in two steps, the embryos were treated with weak Curis solution (1% (v/v) hydrogen peroxide [Merck, Cat#1072090500], 0.5x SSC and 5% formamide) at 4°C in the dark overnight. Finally, the embryos were washed twice in PBS before imaging them in PBS on a thick agarose dish by light microscopy.

# **Processing of External Datasets**

High-time (30-min intervals) resolution of total and poly(A) RNA-Seq (GSE65785) was processed as reported in the original publication (Owens et al., 2016). In addition, intron read counts were corrected by spike RNA-derived normalization factors. For visualization normalized exon and intron counts were scaled to the maximal count detected across the time course and fitted using cubic smoothing splines from 0 to 23.5 hpf: smooth.spline(1:48, x, spar=0.6). Other RNA-Seq (GSE81458) and ChIP-Seq (GSE67974, GSE30146, GSE53654 and GSE72657) were processed as described in detail above except for H3K4me3 and H3K36me3 whose enriched regions were detected as follows: findPeaks -style histone - fragLength 175 -inputFragLength 175 -fdr 0.001 -gsize 1.435e9 -F 2 -C 1 -region -size 350 - minDist 500. Thus, we detected significant regions of histone modifications (-style histone) of at least the lengths of two DNA fragments (-size 350) and being separated by at least 500 bp from each other.

# **Generation of Plots and Heatmaps**

Genomic snapshots were generated with the IGV genome browser v2.4-rc6 (Robinson et al., 2011). All plots and heatmaps were generated using R v3.5.1 (http://cran.r-project.org/). The following add-on R and Bioconductor packages were used for sorting and graphical visualization of data: alluvial v0.1-2 (Michał Bojanowsk), beeswarm v0.2.3 (Aron Eklund), circlize v0.4.5 (Gu et al., 2014), complexHeatmap v1.20.0 (Gu et al., 2016a), dplyr v0.7.8, ggplot2 v3.1.0 (Wickham, 2016), gplots v3.0.1 (Gregory Warnes and colleagues), GenomicFeatures v1.34.1 (Lawrence et al., 2013), GenomicRanges v1.38.0 (Lawrence et al., 2013), HilbertCurve v1.12.0 (Gu et al., 2016b), limma v3.38.2 (Ritchie et al., 2015), rtracklayer 1.42.1 (Lawrence et al., 2009) and seriation v1.2-3 (Hahsler et al., 2008).

## **QUANTIFICATION AND STATISTICAL ANALYSIS**

No statistical method was used for determining sample size; rather, we followed the literature to select the appropriate sample size. The experiments were not randomized. Due to the nature of experiments, the authors were not blinded to group allocation during data collection and

analysis. Only viable embryos were included in the analysis. Frequencies of shown morphological phenotypes and WMISH patterns are included in every image. The significance of over-represented GO terms was based on hypergeometric tests. Significances of non-normally distributed data points (gene features) across ZGA were calculated using paired Wilcoxon rank-sum tests (alternative hypothesis 'less'). The effect size ( $r_{effect}$ ) was estimated from the standard normal deviate of the Wilcoxon p-value (p) as previously described (Rosenthal, 1991),  $r_{effect}$ =Z/sqrt(N), where Z=qnorm(1-p/2) is the standardized Z-score and N is the number of observations.

For RNA-Seq, biological triplicates were used to account for transcriptional variability between clutches. Each LOF experiment has its own control embryos collected in parallel from the same mothers: exp. #1 (α-amanitin), uninjected embryos; exp. #2 (BMP or Nodal LOF), DMSO-treated embryos; exp. #3 (Wnt LOF), uninjected embryos; exp. #4 (mPou5f3/Sox3 LOF), uninjected embryos; exp. #5 (mVegT LOF), uninjected embryos; exp. #6 (single and combinatorial LOFs of Wnt, Nodal and BMP), DMSO-treated embryos; exp. #7 (combinatorial LOF of 4 zygotic T-box TFs), control MO-injected embryos. The gene expressions of control MO-injected embryos of exp. #2 and #5 were normalized to their corresponding uninjected embryos. The mean of these normalizations and conservative FDR estimations (i.e., higher FDR of the two likelihood ratio tests) were used for the comparison with LOF conditions. RNA-Seq libraries from each experiment were generated simultaneously to mitigate any batch effects. The FDR was controlled for multiple comparisons according to the Benjamini-Hochberg procedure. The exact computational implementation of differential expression analysis is outlined on GitHub (see below).

## **DATA AND SOFTWARE AVAILABILITY**

Sequencing reads (FASTQ files) and raw RNA-Seq read counts reported in this paper are available in the GEO database (www.ncbi.nlm.nih.gov/geo) under the accession numbers GSE113186 and GSE122551. All analyses were performed in R v3.5.1 (Bioconductor v3.8), Perl v5.18.2 (https://www.perl.org) and Python v2.7.12 (http://www.python.org) as detailed above. The R code, genome annotation, intermediate datasets and graphs are available on GitHub at <a href="https://github.com/gegentsch/SpatioTemporalControlZGA">https://github.com/gegentsch/SpatioTemporalControlZGA</a>. Original datasets are also available on Mendeley Data at <a href="https://dx.doi.org/10.17632/jn466b4n8v.1">https://dx.doi.org/10.17632/jn466b4n8v.1</a>.

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