

# The Earliest Transcribed Zygotic Genes Are Short, Newly Evolved, and Different across Species

Patricia Heyn,<sup>1</sup> Martin Kircher,<sup>2,4</sup> Andreas Dahl,<sup>3</sup> Janet Kelso,<sup>2</sup> Pavel Tomancak,<sup>1,\*</sup> Alex T. Kalinka,<sup>1,6</sup> and Karla M. Neugebauer<sup>1,5,\*</sup>

<sup>1</sup>Max Planck Institute of Molecular Cell Biology and Genetics, Pfotenhauerstrasse 108, 01307 Dresden, Germany

<sup>2</sup>Max Planck Institute for Evolutionary Anthropology, Deutscher Platz 6, 04103 Leipzig, Germany

<sup>3</sup>Deep Sequencing Group SFB 655, DFG Research Center for Regenerative Therapies (CRTD), Biotechnology Center (Biotec), Technische Universität Dresden, Fetscherstrasse 105, 01307 Dresden, Germany

<sup>4</sup>Department of Genome Sciences, University of Washington, 3720 15th Avenue NE, Seattle, WA 98195, USA

<sup>5</sup>Present address: Molecular Biophysics and Biochemistry, Yale University, 333 Cedar St, New Haven, CT 06520, USA

<sup>6</sup>Present address: Institute of Population Genetics, Vetmeduni Vienna, Veterinärplatz 1, 1210 Vienna, Austria

\*Correspondence: [tomancak@mpi-cbg.de](mailto:tomancak@mpi-cbg.de) (P.T.), [karla.neugebauer@yale.edu](mailto:karla.neugebauer@yale.edu) (K.M.N.)

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## SUMMARY

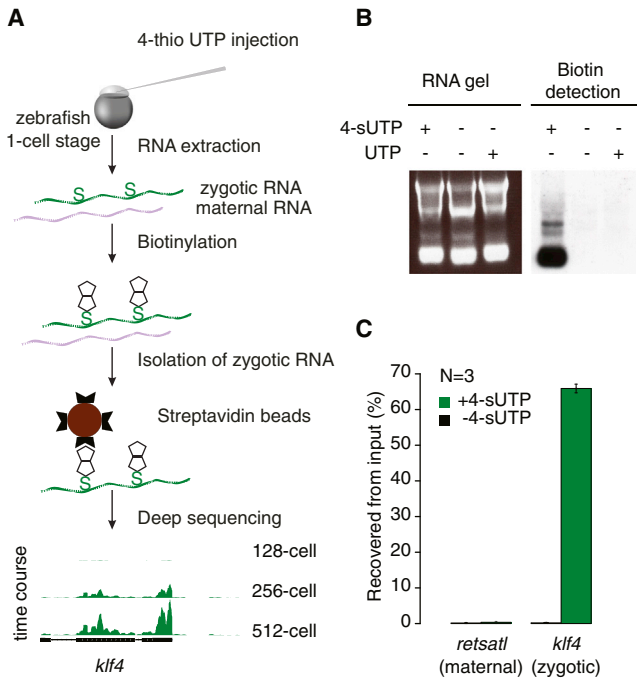
The transition from maternal to zygotic control is fundamental to the life cycle of all multicellular organisms. It is widely believed that genomes are transcriptionally inactive from fertilization until zygotic genome activation (ZGA). Thus, the earliest genes expressed probably support the rapid cell divisions that precede morphogenesis and, if so, might be evolutionarily conserved. Here, we identify the earliest zygotic transcripts in the zebrafish, *Danio rerio*, through metabolic labeling and purification of RNA from staged embryos. Surprisingly, the mitochondrial genome was highly active from the one-cell stage onwards, showing that significant transcriptional activity exists at fertilization. We show that 592 nuclear genes become active when cell cycles are still only 15 min long, confining expression to relatively short genes. Furthermore, these zygotic genes are evolutionarily younger than those expressed at other developmental stages. Comparison of fish, fly, and mouse data revealed different sets of genes expressed at ZGA. This species specificity uncovers an evolutionary plasticity in early embryogenesis that probably confers substantial adaptive potential.

## INTRODUCTION

In all metazoans, the fertilized embryo is provided with proteins and RNAs deposited by the mother during oogenesis. These maternal stores support early embryonic cell divisions before the onset of transcription at zygotic genome activation (ZGA), which occurs after a stereotypical number of cell cycles (Baroux et al., 2008; Tadros and Lipshitz, 2009). The mechanisms regu-

lating ZGA are not fully understood in any species, though recent advances in *Drosophila melanogaster* and *Danio rerio* have begun to identify the maternally provided transcription factors governing the onset of at least some zygotically activated genes (Baroux et al., 2008; De Renzi et al., 2007; Lee et al., 2013; Leichsenring et al., 2013; Tadros and Lipshitz, 2009). In the zebrafish, these studies have focused on a period of robust zygotic transcription that follows the midblastula transition (MBT) when cell cycles begin to lengthen (Kane and Kimmel, 1993). However, some genes are transcribed much earlier. For example, zebrafish MBT occurs at the 1,000-cell stage (1K), yet several studies indicate that first gene transcription might occur as early as the 64-cell stage (Giraldez et al., 2006; Lindeman et al., 2011; Mathavan et al., 2005; O'Boyle et al., 2007). These first zygotic transcripts may play crucial roles in the overall process of ZGA, which continues for several hours during the period of rapid cell division that precedes morphogenesis. Therefore, determining the precise timing and identity of the earliest genes transcribed is central for understanding developmental mechanisms.

The gene expression machinery is confronted by a unique set of challenges during embryogenesis, because the production of functional RNA products is limited by the time it takes to transcribe and process RNA (Swinburne and Silver, 2008). Because transcription shuts down at mitosis and incomplete transcripts are “aborted,” short early embryonic cell cycles are thought to impose a time limit on the genes that can be expressed (Shermoen and O'Farrell, 1991; Tadros and Lipshitz, 2009). The notion that cell-cycle length restricts the potential for gene expression to short genes received support from seminal work in fly development (McHale et al., 2011; McKnight and Miller, 1976; Rothe et al., 1992). Thus, we would assume that early embryos may only be able to express short genes with relatively simple exon-intron architecture. Alternatively, early embryos may have evolved mechanisms for overriding transcript abortion; if so, expression of longer genes might occur. Indeed, one study in zebrafish did identify long genes with many introns among the earliest expressed genes (Mathavan et al., 2005). Thus,



**Figure 1. Metabolic Labeling with 4-sUTP Identifies Zygotically Transcribed Genes**

(A) Schematic of the method for positive selection of zygotic transcripts. Embryos are microinjected with 4-sUTP and grown to the desired developmental stage. Total RNA is extracted and biotinylated in a thiol-specific manner. Biotinylated RNA is captured on magnetic streptavidin beads, extensively washed, and used for downstream applications; here, deep sequencing at three developmental stages and the outcome showing gene coverage for *klf4* is depicted.

(B) Northern blot from 4-sUTP-injected, UTP-injected, or uninjected wild-type control embryos (3 hr postfertilization [hpf]). The panel on the right shows “Northwestern” detection of the biotinylated 4-sUTP incorporated in the RNA.

(C) Quantification of RNA isolation efficiency by qRT-PCR. Plotted are average values ( $\pm$ SD) for percent recovered from input for one maternal gene and one zygotic gene from 3hpf 4-sUTP (+4-sUTP)-microinjected and wild-type control embryos (–4-sUTP) ( $n = 3$ ).

See also Figures S1 and S2.

comprehensive identification of transcribed genes in early embryos provides an opportunity to interrogate cellular and developmental mechanisms.

We sought to address early zygotic transcription in the zebrafish, where the full complement of genes transcribed at the very beginning of ZGA remains unknown. Several studies have performed total steady-state RNA sequencing (RNA-seq) from early embryonic samples (Anes et al., 2011; Pauli et al., 2012; Vesterlund et al., 2011). However, uncovering which zygotic genes are transcribed earliest is hampered by the presence of massive amounts of maternally loaded RNA. Not only are newly transcribed RNAs relatively less abundant, but also “maternal-zygotic” RNAs—those that are maternally provided and are additionally synthesized at ZGA—cannot be deduced from total pools (Supplemental Discussion). To overcome this, recent SNP analysis of maternal and paternal genomes was used to demonstrate the zygotic expression of 3,342 protein-coding genes from

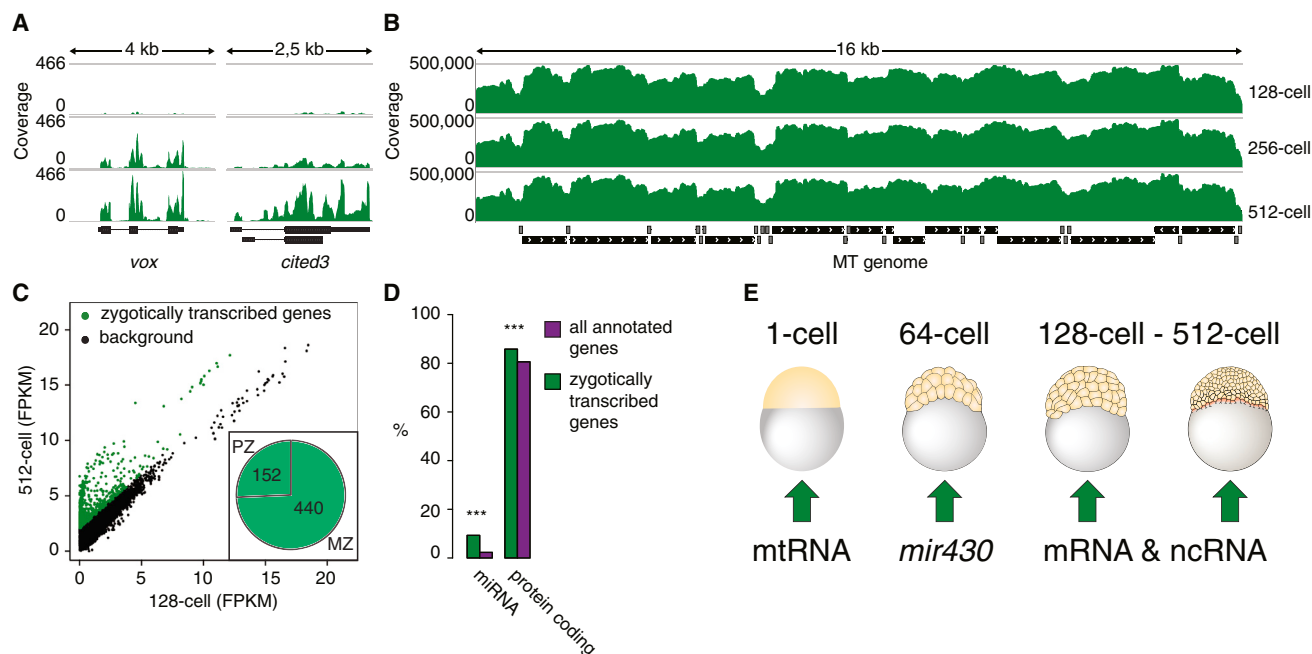
MBT onwards (Harvey et al., 2013). However, SNP analysis is not a comprehensive method, because only  $\sim$ 25% of all genes contained diagnostic SNPs. Previously, other specialized methods have been developed to identify zygotically transcribed genes in flies and plants (De Renzis et al., 2007; Lott et al., 2011; Nodine and Bartel, 2012). Here, we identify the earliest zygotic transcripts in the zebrafish through the establishment of a method to isolate newly transcribed RNA by metabolic labeling of RNA from staged embryos. This is a comprehensive method that allows us to extend robust analysis to vertebrates. We used this data set to compare the first zygotically transcribed genes in fish, fly, and mouse and address whether the set of genes expressed at ZGA is evolutionarily conserved. The results shed light on both the regulation of transcription onset and the evolution of early animal development.

## RESULTS AND DISCUSSION

In order to identify the full complement of the earliest zygotic transcripts, we established a protocol based on RNA metabolic labeling and positive selection of newly synthesized transcripts (Figure 1A; Figure S1A). Contamination from maternal RNA was minimal (Figures 1B and 1C; Figures S1B and S1C), and recovery of zygotic transcripts (e.g., the known zygotic gene *klf4*) was highly efficient (Figure 1C). Therefore, embryos injected at the one-cell stage with 4-thio-UTP (4-sUTP) were allowed to develop to the 128-, 256-, and 512-cell stages before extraction, biotinylation, purification of labeled RNA, and deep sequencing (Figure 1A). We focused on this early sequence of cell divisions because several genes are known to be transcribed before MBT and active chromatin marks have been detected at the 256-cell stage (Lindeman et al., 2011; Mathavan et al., 2005). Coverage of many detected transcripts, such as *klf4*, *vox*, and *cited3*, increased during the cell cycles examined (Figures 1A and 2A), indicating that transcription begins during this time window. Therefore, accumulation of metabolically labeled transcripts over time was used as a signature for robust identification of early zygotic transcripts.

Although it is believed that zygotic genomes are transcriptionally silent before ZGA, RNAs transcribed from the mitochondrial genome were abundant at all three developmental time points (Figure 2B; Figure S2). These data suggest that mitochondrial RNA polymerase is active before the 128-cell stage and that expression has reached steady-state levels. Indeed, independent quantitative RT-PCR (qRT-PCR) experiments show that mitochondrial RNA precursors are clearly detectable from fertilization onward (Figures S3A–S3F). Constitutive mitochondrial transcriptional activity is likely related to the fact that the mitochondrial genome is not packaged into chromatin or subject to DNA methylation (Holt et al., 2007; Potok et al., 2013). Therefore, mitochondrial genome activity is independent of nuclear genome regulation, and the dogma that the early zygote is transcriptionally silent (Baroux et al., 2008; De Renzis et al., 2007; Kane and Kimmel, 1993; Lee et al., 2013; Leichsenring et al., 2013; O’Boyle et al., 2007; Tadros and Lipshitz, 2009) is not strictly correct.

Nuclear genome activity was detected between the 128- and 512-cell stages, with clear increases in transcript abundance observed for 592 nuclear genes, comprising 670 transcript



**Figure 2. Timing of Protein-Coding, miRNA, and Mitochondrial Gene Transcription during ZGA**

(A) Example of increasing read coverage over the zygotically transcribed genes (*vox* and *cited3*). The architecture of the respective genes is displayed below. (B) Display of constant high read coverage over the mitochondrial genome. The architecture of the zebrafish mitochondrial genome is displayed below. Protein-coding genes are colored in black and tRNAs in gray.

(C) Scatterplot of abundance for zygotically transcribed genes (green,  $n = 592$ ). Genes in black are unchanged between the developmental time points. To avoid negative values in the log<sub>2</sub> scale, 1 was added to all FPKM values before transformation. The inset shows the proportion of purely zygotically transcribed genes (PZ) and maternal-zygotic expressed genes (MZ).

(D) Identification of overrepresented transcript types in zygotically transcribed genes compared to the overall distribution in all annotated zebrafish genes. The proportions of each biotype among the zygotically transcribed genes (green bars) and among all annotated genes (purple) are plotted in percent. The stars indicate significant enrichment over random distribution (hypergeometric test, \*\*\* equals  $p$  value  $\leq 0.001$ ; Table S3). For other biotypes present, see Figure S5A.

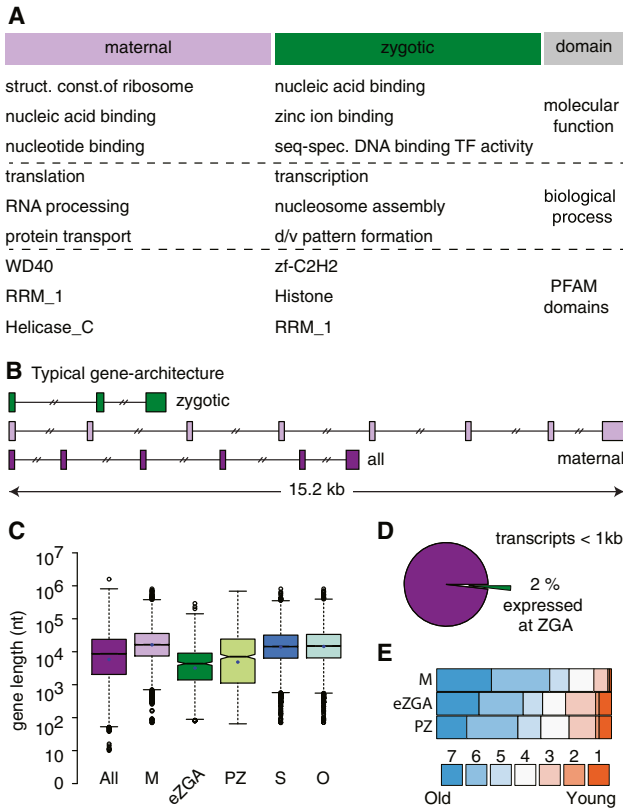
(E) Schematic showing developmental stages and the detection of different types of newly transcribed genes. See also Figures S3–S5 and Tables S1, S2, and S3.

isoforms (Figure 2C; Tables S1 and S2; Figures S2B and S4A–S4C). Transcription inhibition by  $\alpha$ -amanitin showed that purification of metabolically labeled RNA is due to incorporation of 4-sUTP during transcription (Figure S4D). Because our RNA-seq data were normalized to constant mitochondrial RNA levels, the identified changes are specific for each gene and not the result of global shifts in RNA populations. Comparison of this data set with the recent study based on SNP analysis (Harvey et al., 2013) revealed that our method detects 350 zygotically transcribed genes that lack informative SNP loci. In addition, 84 SNP-containing genes overlapped with our data set. Interestingly, Pou5f1 was recently identified as a major regulator of ZGA in zebrafish embryos (Lee et al., 2013; Leichsenring et al., 2013). We find enrichment of Pou5f1 and Sox2 peaks among ~50% of our genes (data not shown), making it plausible that Pou5f1 drives expression of at least some of these early zygotic genes.

The zygotically expressed genes identified in our study primarily encode proteins and microRNAs (miRNAs) transcribed by RNA polymerase II (Figure 2D; Figures S5A and S5B; Table S3). As expected, we detected *mir430*, which has demonstrated roles in embryonic RNA regulation (Giraldez et al., 2006), as well as *mir19a*, a newly identified zygotic transcript (Figure S4C). In

addition, noncoding RNAs transcribed by Pol III were robustly identified (Table S1). Of the 592 genes detected, 152 were “purely zygotic” (PZ) and not detected in the maternal pool (Figure 2C, inset). Increased detectability of RNA precursors over the time course is consistent with de novo transcription (Figure 2A; Figures S4A–S4C). Note that many introns from robustly transcribed genes are poorly covered (see Figure 2A), likely because pre-mRNA splicing is a fast cotranscriptional process (Brugiolo et al., 2013). We conclude that early ZGA begins with transcription of *mir430* genes by the 64-cell stage, followed by nuclear protein-coding genes and other noncoding RNAs at the 256- to 512-cell stage (Figures 2E and S4C). The remaining 74% (440) of genes were “maternal-zygotic” (MZ; Figure 2C, inset), meaning that transcripts were maternally provided and then synthesized again at early ZGA. MZ genes account for only 3% of the total number of maternally expressed genes (14,500) previously identified (Pauli et al., 2012). This suggests that the re-expression of selected maternal genes at early ZGA may be specifically required.

Comparison of maternal and zygotic protein-coding genes reveals dramatic differences in function, reflecting a trend toward RNA regulation for maternal transcripts and DNA regulation



**Figure 3. Zgotically Transcribed Genes Are Short and Intron Poor**

(A) Gene Ontology term and protein families (PFAM) domain analysis of maternally expressed and zgotically transcribed genes. The three most enriched terms or domains are shown.

(B) Stick diagram of typical gene architecture for zgotic, maternal, and all annotated zebrafish transcripts. Drawn to scale is the median length of the genes (zgotically transcribed median = 3,874 bp; maternally expressed median = 15,202 bp and all annotated genes median = 8,644bp) and the first and last exons. For internal exons, the population median for all exons per transcript is drawn. Introns are not to scale; median numbers of introns (zgotic: 2, maternal: 7; and all: 5) are shown.

(C) Distribution and median gene length for different developmental gene categories for *D. rerio*. The categories correspond to all annotated genes (all), maternal genes (M), early zgotic genes (eZGA), purely zgotic genes expressed after MBT (PZ), genes expressed at somitogenesis (S), and genes expressed at organogenesis (O). For p values of all pair-wise comparisons, see Table S6. Table S11 lists all data sets used.

(D) Pie chart showing the proportion of zgotic transcripts that are shorter than 1kb among all (n = 5,822).

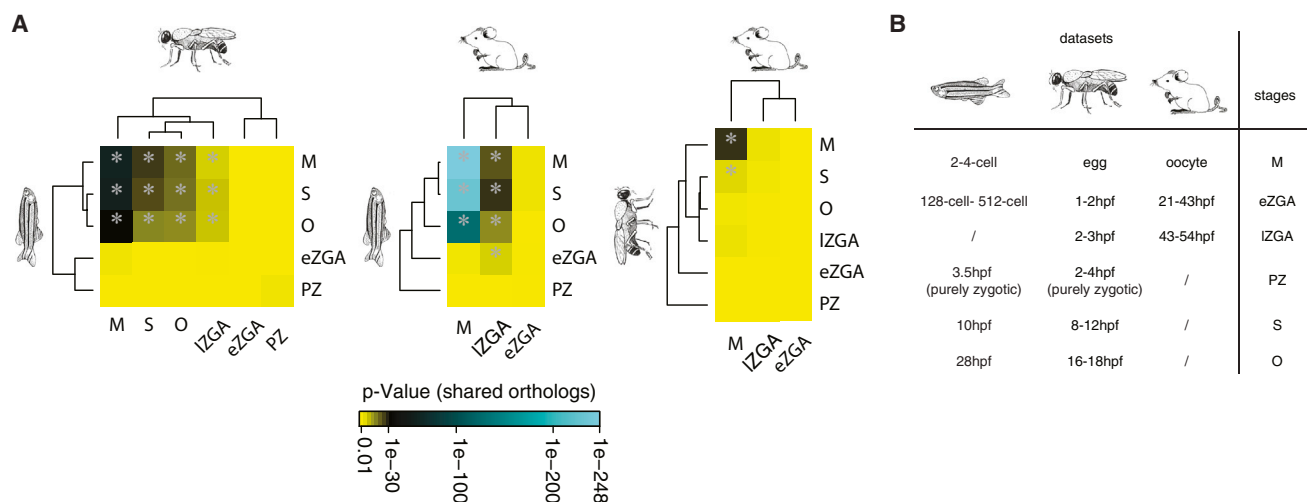
(E) Stacked bar graphs showing the normalized distribution of the inferred age of zebrafish genes expressed at maternal stage (M) and at early ZGA (eZGA) and purely zgotic genes expressed after MBT (PZ). Gene age is retrieved from Ensembl protein trees depicted schematically in Figure S8C. Different colors correspond to numbers depicted underneath the bar graph and indicate the node (clade) in the protein tree in which the genes emerged, where smaller numbers represent more recent emergence. See also Figures S5–S8 and Tables S4, S5, S6, and S7.

for zgotic transcripts (Figure 3A; Tables S4 and S5). For example, ribosomes and spliceosomes are abundantly contributed maternally to facilitate high levels of translation and splicing during early embryogenesis (Amsterdam et al., 2004; Strzelecka

et al., 2010). Interestingly, 143 of 144 genes annotated for “structural constituent of the ribosome” are detected among maternal transcripts, while only 13 of these genes were detected among zgotic transcripts. Because zgotic rRNA and small nuclear RNA expression were also not detected (Figure S5B), early embryos may rely exclusively on maternally provided ribosomes and spliceosomes. In contrast, zgotic RNAs encode DNA binding proteins, histones and histone variants, and chromatin modifiers (Figure 3A; Tables S4 and S5). A number of transcription factors with later roles in patterning and organogenesis, such as Forkhead and Dickkopf proteins, are among the purely zgotic transcripts; roles for these factors prior to the first steps of morphogenesis are thus far unknown (Sprague et al., 2003). On the other hand, the observed early zgotic expression of cell-cycle regulators, such as *c-fos* (*fos*) and cyclin-dependent kinase inhibitor (*cdkn1a*), could reflect their involvement in establishing gap phases in the cell cycle at MBT.

Gene transcription and RNA processing take time, and transcription elongation is usually aborted at mitosis. Cell-cycle lengths in the zebrafish are only 15 min long prior to MBT, suggesting that zgotic gene lengths may also be limited. However, work in zebrafish suggested that very long genes can indeed be expressed at early time points (Mathavan et al., 2005). Analysis of the architecture of our 592 zgotically expressed genes shows that early zgotic genes are four times shorter than maternal genes (Figure 3B and 3C; Figures S5C and S5D; Tables S6 and S7). They are relatively intron poor or intronless (Figure 3B; Figures S5E and S5F). These findings suggest that many of the long genes identified in the previous study might reflect contaminating levels of maternal RNA. This highlights the utility of a positive selection method, such as the one presented here, to identify all classes of newly transcribed genes. Orthologs of the zebrafish early zgotic genes in the very compact *Fugu* as well as the enlarged *Coelacanth* genomes are also short (Figure S6; Table S6). To address whether gene shortness is specific for ZGA or might coincidentally reflect the types of genes enriched in the zgotic pool, the gene architecture of all transcription factors was analyzed. We show that transcription factor genes are not short in general, yet those that are zgotically expressed are significantly shorter (Figure S7A; Table S6; Mann-Whitney *U* test,  $p < 0.001$ ). Furthermore, if transcription factors are eliminated from the zgotic pool, we still observe an enrichment of short genes (Figure S7B). In contrast, maternally expressed genes as well as genes expressed during differentiation and organogenesis (Aanes et al., 2011; Pauli et al., 2012) are significantly longer than the genome-wide median (Figures 3B and 3C; Figures S5C, S5D, S7A; Tables S6 and S7; Mann-Whitney *U* test,  $p < 0.001$ ). Therefore, zgotic genes are short, and maternal genes are long.

The shortness of early zgotic genes in fruit fly (Figure S6B) and mosquito has been taken as evidence that cell-cycle length may determine gene expression in early embryos (Biedler et al., 2012; De Renzis et al., 2007; McKnight and Miller, 1976; Rothe et al., 1992). Moreover, one model proposes that gene length actively determines which genes are expressed at MBT; in this model, transcription activation is widespread but transcript abortion at mitosis prevents long genes from being expressed (Swinburne and Silver, 2008; Tadros and Lipshitz, 2009). We



#### Figure 4. Expression of Early Zygotic Genes Is Evolutionarily Divergent

(A) Clustered heatmaps for pairwise ortholog comparisons between species (fly, fish, and mouse) at and between respective developmental stages. See [Table S11](#) for a list of the data sets used. Depicted are color-coded p values describing the significance of shared ortholog enrichment between species. Enrichments were considered significant with a p value smaller than the Bonferroni cutoff of  $0.05/63 = 0.00079$ , indicated by gray asterisks. Hierarchical clustering is solely for visualization purposes.

(B) Overview of developmental stages used for analysis of shared orthologs in the three species (fly, fish, and mouse). See also [Figure S9](#) and [Tables S8, S9, S10, and S11](#).

tested this hypothesis by examining all genes less than 1 kb long (less than one-third the length of observed zygotic genes) in order to objectively evaluate expression of short genes. Only 2% of >5,000 short genes are expressed zygotically in zebrafish ([Figure 3D](#)). Therefore, our data rule out the possibility that zygotic transcription is determined by short gene length alone. Interestingly, the presence of introns in many of these zygotic genes, though short, may be an important factor in gene activation; it was recently shown that intron content positively regulates transcriptional output and quality ([Bieberstein et al., 2012](#)).

Short genes tend to be evolutionarily young ([Grzybowska, 2012](#); [Neme and Tautz, 2013](#); [Shabalina et al., 2010](#)), suggesting that early zygotic genes may have evolved recently. To test this, we compared the evolutionary age of genes expressed at different stages of zebrafish development. Genes expressed at ZGA were indeed younger than genes expressed maternally and later during tissue specification ([Figure 3E](#); [Figures S8A–S8D](#)). Extending this analysis to published fruit fly data ([De Renzis et al., 2007](#); [Graveley et al., 2011](#); [Hamatani et al., 2004](#); [Xue et al., 2013](#); [Zeng et al., 2004](#)) ([Figures S8A and S8C](#)), an even tighter correlation regarding gene age was revealed ([Figure S8A](#)). The results show that genes expressed at early ZGA in the fly and the fish are significantly younger than expected ([Figure S8D](#);  $\chi^2$  test,  $p < 2.22 \times 10^{-16}$ ). This was not the case in the mouse ([Figure S8](#)). Mice and fish are both vertebrates and share a common ancestor more recently than they do with flies. However, fish and mice differ in the dynamics of early embryogenesis; for example, early cell division times differ by almost two orders of magnitude. Interestingly, cellular dynamics are more similar between fish and fly embryos, where cell-cycle lengths are only 8–15 min before ZGA. Therefore, cellular dynamics may be a major determinant of gene expression at ZGA across phyla and species.

Fly, fish, and mouse early embryos clearly differ in many ways. For example, in the syncytial blastoderm of the fly mitosis takes place without cytokinesis, while partial cleavages of the egg take place in fish; cell-cycle lengths and patterning strategies also differ. However as ZGA precedes cellular differentiation in all metazoans, one might expect specific mechanisms to have evolved in the last common ancestor and to have been retained in the animals we study today. The observation that early zygotic genes tend to be evolutionarily younger, while maternal genes are more ancient, prompted us to ask whether the sets of genes expressed in the early embryo at different time points are evolutionarily conserved. We conducted a comparative study using our data set and published data on fruit fly and mouse ([De Renzis et al., 2007](#); [Hamatani et al., 2004](#); [Xue et al., 2013](#)) to specifically interrogate the sets of genes in different species for shared expression of orthologs. This analysis reveals a strong enrichment of shared orthologs among maternally provided genes ([Figure 4](#); [Figure S9A](#); [Tables S8 and S9](#); hypergeometric test,  $p < 0.00079$ ). Between fruit fly and zebrafish, 2,773 shared orthologs were identified among the maternal genes, representing 19.1% of the zebrafish maternal gene pool ([Table S9](#)). The same trend was observed among the maternal genes of fruit fly and mouse ([Figure 4](#); [Figure S9B](#); [Table S8](#)). Maternal genes are also highly enriched for essential genes in all three species ([Table S10](#); one-tailed hypergeometric test,  $p < 0.005$ ). These data indicate that maternally contributed RNAs and proteins, and the essential functions they provide, constitute evolutionarily conserved features of early embryogenesis.

If early zygotic gene products also play a role in the awakening of the rest of the genome, then we would expect to reveal shared orthologs among the early zygotic genes in flies, fish, and mice as well. The recent identification of so-called hub genes, shared in

expression between mouse and human embryos, supports this notion (Xue et al., 2013). However, the identification of shared orthologs expressed among the three zygotic gene pools yielded a poor level of overlap that would be expected by chance (Figure 4; Figures S9A–S9C; Table S8; hypergeometric test,  $p = 1$  [fly–fish early ZGA],  $p = 0.1027935$  [fish–mouse early ZGA],  $p = 1$  [fly–mouse early ZGA]). Note that our analysis only considers orthologs that are expressed in both species at specific developmental stages (shared orthologs). In contrast to the zygotic genes, adult traits associated with similar small numbers of genes showed strong enrichment across the same species (Figure S9D; Table S8). Remarkably, there are no shared orthologous early zygotic genes expressed in both fruit fly and zebrafish. Moreover, purely zygotic genes were depleted of essential genes (Table S10), consistent with the observation that young genes are less likely to be essential (Chen et al., 2012). Therefore, in stark contrast to maternal genes, early zygotic genes are completely different between arthropods and vertebrates as well as among vertebrates, suggesting species-specific functions.

Many classes of developmental events, such as signaling by growth factors, establishment of cell polarity, and the execution of differentiation programs, show a high degree of evolutionary conservation in the genes and gene products involved. On the other hand, mechanisms underlying other key events like sex determination and dosage compensation differ widely. We show that maternal transcripts are long, evolutionarily older, and tend to be shared across distantly related species. This suggests that the complement of maternal RNA and protein provided during oogenesis is critically important for the regulation of early embryogenesis by primarily RNA-driven mechanisms such as translation. In contrast, early zygotic genes are short, evolutionarily younger than other developmental genes, and not orthologous to early zygotic genes in other species. This indicates that ZGA occurs during a remarkably flexible period of development in which the order of gene activation is likely sculpted by species- or lineage-specific cellular mechanisms, such as morphology or the usage of yolk or trophoderm to support the zygote (Kalinka and Tomancak, 2012). Such a correspondence between early, species-specific aspects of embryogenesis and the species specificity of the underlying molecular components of ZGA is consistent with the hourglass model of developmental evolution, in which the nature of early development reflects the diverse reproductive and ecological strategies of individual species (Duboule, 1994; Raff, 1996). However, the expression of shared sets of orthologs at the maternal stage does not necessarily imply that these components of early development are being used in identical ways in these different species. Rather, our results suggest that the unique constraints acting during ZGA provide a window of opportunity for the expression of evolutionarily younger, short genes that are capable of adding new functions to the zygotic gene expression program.

## EXPERIMENTAL PROCEDURES

### Metabolic Labeling

Fertilized zebrafish embryos were obtained according to approved protocols at the MPI-CBG zebrafish facility. Embryos were injected at the one-cell stage

with 1 nl 50 mM 4-sUTP (Ambion, Trilink) and collected at the desired stage. Biotinylation and purification of labeled RNA was performed as described previously (Zeiner et al., 2008) with minor modifications.

### RNA-Seq Analysis

Isolated RNA was amplified with the WT-Ovation One-Direct RNA Amplification System (NuGEN), converted into TruSeq sequencing libraries (Illumina), and sequenced on a HiSeq2000 instrument (Illumina). For a list of primers, see Table S12. Raw reads were trimmed and mapped with TopHat 1.3.3 (Trapnell et al., 2009) to the zebrafish genome assembly Zv9/GCA\_000002035.2. Quantification of gene/transcript expression was performed with Cufflinks 2.0.2 (Trapnell et al., 2010). To normalize between different time points in our study, the sum of FPKM values for protein-coding mitochondrial transcripts/genes was assumed to be constant.

### Gene Age and Ortholog Analysis

The age of each gene was determined by the age of the oldest node in its protein phylogeny (Piasecka et al., 2013) using protein trees. Protein trees, generated by EnsemblCompara (Vilella et al., 2009), a phylogeny-aware, gene-tree-building pipeline based on BLASTP, were downloaded from Ensembl for mouse and fish, and from Ensembl Metazoa for the fly. Enrichment of shared orthologs was tested for each pair of species and for genes expressed at several developmental stages. One to two orthologs were allowed for fish comparisons due to the whole-genome duplication in teleosts (1:2, fly:fish, mouse:fish), and a variant of the hypergeometric test was applied to test for enrichment (Kalinka, 2013). Enrichments were considered significant with a  $p$  value smaller than the Bonferroni cutoff of  $0.05/63 = 0.00079$ .

### ACCESSION NUMBERS

Raw and processed RNA-seq data sets have been deposited into NCBI Gene Expression Omnibus (<http://www.ncbi.nlm.nih.gov/geo/>) under accession number GSE47709.

### SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Discussion, Supplemental Experimental Procedures, nine figures, and twelve tables and can be found with this article online at <http://dx.doi.org/10.1016/j.celrep.2013.12.030>.

### AUTHOR CONTRIBUTIONS

P.H. and K.M.N. designed the study; P.H. performed all experiments; A.D. provided expertise for library preparation; M.K. and J.K. developed and implemented analysis tools for the primary data; M.K., P.H., and A.T.K. performed secondary analyses; A.T.K. developed methods for ortholog analysis; and P.H. and K.M.N. wrote the manuscript with support from P.T., M.K., J.K., and A.T.K.

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