

RESEARCH REPORT

Histone concentration regulates the cell cycle and transcription in early development

Sudarshan Chari*, Henry Wilky*, Jayalakshmi Govindan and Amanda A. Amodeo†

ABSTRACT

The early embryos of many animals, including flies, fish and frogs, have unusually rapid cell cycles and delayed onset of transcription. These divisions are dependent on maternally supplied RNAs and proteins including histones. Previous work suggests that the pool size of maternally provided histones can alter the timing of zygotic genome activation (ZGA) in frogs and fish. Here, we examine the effects of under- and overexpression of maternal histones in *Drosophila* embryogenesis. Decreasing histone concentration advances zygotic transcription, cell cycle elongation, Chk1 activation and gastrulation. Conversely, increasing histone concentration delays transcription and results in an additional nuclear cycle before gastrulation. Numerous zygotic transcripts are sensitive to histone concentration, and the promoters of histone-sensitive genes are associated with specific chromatin features linked to increased histone turnover. These include enrichment of the pioneer transcription factor Zelda, and lack of SIN3A and associated histone deacetylases. Our findings uncover a crucial regulatory role for histone concentrations in ZGA of *Drosophila*.

KEY WORDS: Chromatin, Maternal to zygotic transition, MZT, Mid-blastula transition, MBT, Zygotic genome activation, ZGA, Nuclear cytoplasmic ratio, Cell cycle

INTRODUCTION

Upon fertilization the embryo must produce enough cells to pattern a functioning organism. Many species accomplish this by undergoing rapid cleavage divisions with little transcription. This is followed by zygotic genome activation (ZGA) (Tadros and Lipshitz, 2009; Vastenhouw et al., 2019; Harrison and Eisen, 2015; Blythe and Wieschaus, 2015a). In many organisms, ZGA coincides with the cell cycle slowing that precedes the onset of morphogenesis during a period called the mid-blastula transition (MBT) (Newport and Kirschner, 1982a,b). In *Drosophila*, the initial divisions are further shortened through omission of cytokinesis, resulting in a syncytium for the first 13 nuclear cycles (NCs), which becomes cellularized when the cycle slows in NC14 (Foe and Alberts, 1983). This switch from transcriptionally silenced, rapid NCs to transcriptionally active, slower NCs is accompanied by changes in the chromatin landscape. In *Drosophila*, transcription factors bind their consensus sequences (Harrison et al., 2010, 2011; Liang et al., 2008; Driever and Nüsslein-Volhard, 1989), RNA polymerase is

gradually recruited to sites of transcription (Gaertner et al., 2012; Chen et al., 2013; Blythe and Wieschaus, 2015b) and nucleosome-free regions open on promoters (Li et al., 2014; Blythe and Wieschaus, 2016). At a larger scale, the genome becomes folded into topologically associated domains, and heterochromatin is differentiated from euchromatin for the first time (Hug et al., 2017; Shermoen et al., 2010; Seller and O'Farrell, 2018).

The early divisions are fueled by maternally supplied RNAs, proteins and metabolites (Newport and Kirschner, 1982a,b; Vastag et al., 2011; Song et al., 2017; Collart et al., 2013; Djabrayan et al., 2019; Liu et al., 2019). Chromatin components, including core histones, are loaded in vast excess of what is required for the first divisions (Horard and Loppin, 2015; Adamson and Woodland, 1974; Woodland and Adamson, 1977; Shindo and Amodeo, 2019). In *Drosophila*, maternal histone stores are sufficient for progression through the MBT, but zygotic histone production is required for progression past the first post-MBT cell cycle (Günesdogan et al., 2010, 2014; Zhang et al., 2018). It has been suggested that overabundant histones compete with transcription factors for DNA binding to repress transcription in the early cycles (Almouzni et al., 1990, 1991; Prioleau et al., 1994; Almouzni and Wolffe, 1995; Amodeo et al., 2015; Joseph et al., 2017). Indeed, pioneer transcription factors are required to evict nucleosomes at ZGA, and altering the concentration of core histones in the early embryos of both *Xenopus* and zebrafish can modulate the timing of ZGA (Liang et al., 2008; Harrison et al., 2010, 2011; Lee et al., 2013; Leichsenring et al., 2013; Amodeo et al., 2015; Joseph et al., 2017). Although the loss of embryonic linker histone H1 has been implicated in timing *Drosophila* ZGA, the role of core histones has not been well characterized (Pérez-Montero et al., 2013).

Here, we examine the effect of histone concentration on the MBT in *Drosophila*. We manipulate maternally deposited histone concentration and examine the effects on cell cycle and transcription. Histone reduction results in slower and fewer nuclear cycles, whereas histone increase results in additional cycles. We demonstrate that cell cycle changes correspond to large changes in zygotic transcription and maternal mRNA degradation consistent with alterations to ZGA. We identify a subset of transcripts as directly histone sensitive. The promoters of histone-sensitive genes are enriched for the pioneer transcription factor Zelda (ZLD) and the absence of the transcriptional repressor SIN3A and associated histone deacetylases (HDACs). Together, these results demonstrate a direct role for histone concentrations in regulating the MBT, provide a list of transcripts that are both directly and indirectly sensitive to histone concentration, and identify the chromatin features that underlie direct histone sensitivity.

RESULTS

Histone concentration regulates the timing of the MBT

To understand the effects of histone concentration on the MBT we reduced maternally supplied histones by downregulating the gene encoding a crucial histone regulator, *Stem-Loop Binding Protein*

Lewis-Sigler Institute for Integrative Genomics, Princeton University, Princeton, NJ 08544, USA.

*These authors contributed equally to this work

†Author for correspondence (aamodeo@princeton.edu)

© A.A.A., 0000-0003-3701-6375

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(*Slbp*) (Sullivan et al., 2001; Dominski et al., 2002; Lanzotti et al., 2002; Iampietro et al., 2014; Lefebvre et al., 2017; He et al., 2018) via maternally driven RNAi (Perkins et al., 2015). Under these conditions, histone H2B was reduced by ~50% and H3 by ~60% at the MBT (Fig. S1A). Approximately 50% of embryos laid by *Slbp* RNAi mothers (henceforth *Slbp* embryos) that form a successful blastoderm do not undergo the final division and attempt gastrulation in NC13 (Fig. 1A, Movie 1). Another ~30% exhibit an intermediate phenotype of partial arrest, with only part of the embryo entering NC14 (Movie 2). A minority of *Slbp* embryos begin gastrulation with all nuclei in NC14. NC12 duration was predictive of NC13 arrest, with NC12 being an average of ~5 min longer in *Slbp* embryos that went on to arrest compared with those that did not arrest ($P=0.0034035$) (Fig. 1C, Table S1, see Materials and Methods).

We first detected cellularization in wild-type (WT) embryos ~20 min into NC14. Partially arrested *Slbp* embryos also began cellularization ~20 min into NC14, with nuclei that arrested in NC13 waiting until the remainder of the embryo had entered NC14 to cellularize. Fully arrested embryos began cellularization ~20 min into NC13, initiating cellularization one cycle early and ~20 min earlier in overall developmental time than WT. Despite their reduced cell number, these embryos form mitotic domains and gastrulate without obvious defects, however they die before hatching (Fig. S2A,B).

To examine the effects of increased histone concentration on developmental timing we monitored cell cycle progression in embryos from *abnormal oocyte* (*abo*) mutant mothers (henceforth *abo* embryos). *abo* is a histone locus-specific transcription factor, the knockdown of which increases the production of replication-coupled histones, particularly H2A and H2B (Berloco et al., 2001). We found that *abo* increased H2B by ~90%, whereas total (combined replication-coupled and replication-independent) H3 was not affected in NC14 embryos (Fig. S1B). Approximately 60% of *abo* embryos displayed fertilization defects or catastrophic early nuclear divisions (Table S1, Tomkiel et al., 1995). Of *abo* embryos that formed a functioning blastoderm, ~6% underwent a complete extra nuclear division before gastrulating in NC15, whereas ~4% underwent a partial extra nuclear division (Fig. 1B, Movies 3,4). Embryos from *abo* mothers that completed total extra divisions had faster NC14s in which they did not cellularize and spent 40–60 min in NC15 before gastrulating (Fig. 1D). This suggests an alteration of the normal transcription-dependent developmental program. In some cases, the cell cycle program and transcriptional program may be decoupled, evidenced by the fact that some *abo* embryos attempted to gastrulate while still in the process of division. *abo* embryos that underwent extra divisions exhibited a range of gastrulation defects including expanded mitotic domains and ectopic furrow formation (Fig. S2A,C, Movies 3,4).

As alterations in histone levels can both decrease and increase the number of divisions before cell cycle slowing, we reasoned that histone levels might affect activation of checkpoint kinase 1 (Chk1, also known as *grp*), which is required for cell cycle slowing at the MBT (Fogarty et al., 1994, 1997; Sibon et al., 1997, 1999). To test this, we crossed a fluorescent biosensor of Chk1 activity into the *Slbp* background (Deneke et al., 2016). We found that even in *Slbp* embryos that did not undergo early gastrulation, Chk1 activity was higher than in WT, consistent with the lengthened cell cycle (Fig. 1E). This result indicates that the observed cell cycle phenotypes in the histone-manipulated embryos are likely mediated through changes in Chk1 activity.

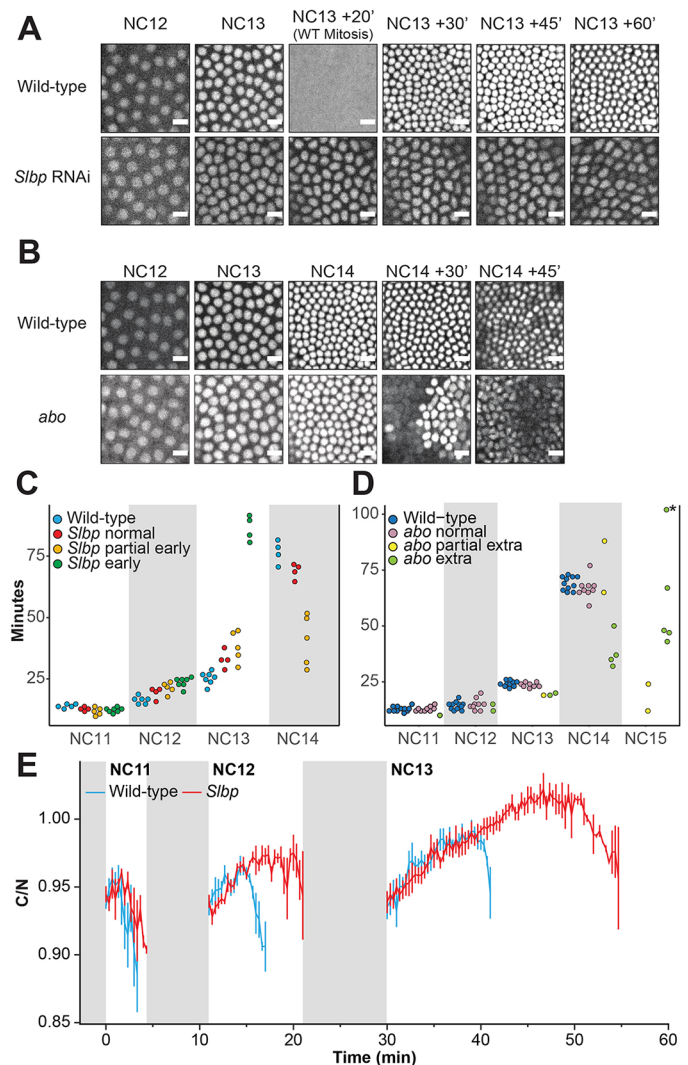


Fig. 1. Changes to replication-dependent histone concentration after cell cycle and timing at the MBT. (A,B) Still frames from time-lapse imaging of wild-type (WT), *Slbp* (histone knockdown, A) and *abo* (histone overexpression, B) embryos from NC12 through gastrulation. Nuclei were visualized using an nls-RFP marker. Approximately 50% of *Slbp* embryos that form a blastoderm do not undergo a 13th mitosis (shown). WT embryos mitose into NC14 ~20 min after entering NC13, but these *Slbp* embryos remain in NC13 where they cellularize, then gastrulate at lower nuclear densities. Approximately 30% of *Slbp* embryos that form a blastoderm undergo a partial 13th mitosis. Conversely, ~6% of observed *abo* embryos have a shortened NC14 before undergoing a full extra mitosis and attempting a catastrophic gastrulation at NC15, and ~4% undergo a partial extra mitosis (shown). Scale bars: 10 μ m. (C) Scatterplot of *Slbp* cell cycle times compared with WT. Cell cycle times were predictive of phenotype with longer early cycles in *Slbp* associated with full arrest in NC13. (D) Scatterplot for *abo* cell cycle times compared with WT. Shorter early cycles in *abo* are associated with extra divisions. NC15 data point with asterisk (*) denotes an embryo that underwent a 16th NC. (E) Cytoplasmic to nuclear ratio (C/N) of the Chk1 biosensor in WT (blue, $n=4$) and *Slbp* (red, $n=4$) embryos from NC11–NC13. Chk1 is prematurely activated in *Slbp* embryos that have lengthened early cell cycles but do not undergo premature gastrulation. Gray boxes represent mitosis. Data are mean \pm s.e.m.

Low histone concentration advances and high histone concentration delays ZGA

As cellularization and gastrulation require zygotic transcription, we suspected that embryos with altered development likely have altered gene expression. We performed single-embryo RNA-seq on staged *Slbp* embryos that remained in NC13 for more than 30 min (Fig. 2A).

We compared these with either NC-matched (NC13) or time-matched (NC14) WT embryos. To control for maternal effects of *Slbp* RNAi, we collected pre-blastoderm stage WT and *Slbp* embryos. We found that the *Slbp* embryos undergo ZGA one NC earlier than WT. We identified ~5000 genes that are differentially expressed (see Materials and Methods for details) between *Slbp* and WT NC13, with ~60% being upregulated (Fig. 2B, Tables S2-S5, S19, S21). The upregulated genes have largely previously been identified as new zygotic transcripts, including cell cycle regulators such as *fruhstart*

(*frs*, also known as *Z600*) and signaling molecules such as *four-jointed* (*ff*), whereas the downregulated genes are enriched for maternally degraded transcripts (Fig. 2C, D, Table S6, S16, S22) (Lott et al., 2011; De Renzis et al., 2007). We believe this represents a coherent change in ZGA timing instead of global transcription dysregulation, as 98% of the genes that are overexpressed in *Slbp* are expressed before the end of NC14 in our control or previously published datasets (Table S20) (Lott et al., 2011). Indeed, the transcriptomes of histone-depleted embryos that stopped in NC13 are more similar to WT NC14 than WT NC13, which suggests a role for cell cycle elongation in ZGA (Fig. S3A, Yuan et al., 2016). Nonetheless, ~1500 genes are differentially expressed between *Slbp* NC13 and WT NC14 without accounting for differences in ploidy. Of these, the majority of the ~1000 overexpressed genes are again associated with zygotic transcription, and downregulated genes associated with maternal products (Fig. S3E, Tables S2-S6, S14, S15). Thus, ZGA is even further accelerated in the histone knockdown than can be explained by purely time alone.

As ZGA is accelerated by histone depletion, we asked whether ZGA would be delayed in the histone overexpression mutant. We performed RNA-seq on pools of *abo* and WT embryos collected 15-30 min into NC14 (Fig. 3A). We identified >1000 genes that are differentially expressed between *abo* and WT, with approximately equal numbers of genes up- and downregulated (Fig. 3B, Fig. S4, Tables S2, S7-S9, S16, S19, S21). As expected, the downregulated genes in *abo* are enriched for previously identified zygotically expressed transcripts (Lott et al., 2011; De Renzis et al., 2007) (Fig. 3D, Tables S6, S14, S16, S22), and upregulated transcripts are enriched for maternally deposited genes. Thus, histone overexpression delays the onset of ZGA.

Zygotically expressed genes, the transcription of which is upregulated by histone depletion and downregulated by histone overexpression, contain many important developmental and cell cycle regulators including: *frs*, *hairy* (*h*), *fushi tarazu* (*ftz*) and *odd-skipped* (*odd*) (Figs 2C, 3C, Tables S4, S7, S8). Conversely, the maternally degraded transcripts that are destabilized by histone depletion and stabilized by histone overexpression include several cell cycle regulators such as *Cyclin B* (*CycB*), *string* (*Stg*, also known as *Cdc25^{string}*) and *Myt1* (Tables S4, S7, S8). Therefore, histone concentration can modulate the expression and stability of specific cell cycle regulators, which may contribute to the onset of MBT.

As histone concentration has previously been implicated in sensing the nuclear-cytoplasmic (N/C) ratio (Amodeo et al., 2015), we compared the genes that are changed in both the histone under- and overexpression embryos with those that had previously been found to be dependent on either the N/C ratio or developmental time (Lu et al., 2009). Both previously identified N/C ratio-dependent and time-dependent genes (Lu et al., 2009) followed the same general trends as the total zygotic genesets (De Renzis et al., 2007), indicating that histone availability cannot explain these previous classifications (Fig. S6, Table S6, S22).

Next, we sought to disentangle the effects of cell cycle length from transcription in the histone overexpression mutant. We performed single-embryo time-course RNA-seq on *abo* and WT embryos collected 3 min before mitosis of NC10-NC13 and 3 min into NC14 (Fig. 3E). In addition, we collected unfertilized embryos (henceforth NC0) of both genotypes to control for differences in maternal contribution. Even with a stringent selection process that accounted for cell cycle time and embryo health (see Materials and Methods, Fig. S5A, B, Tables S2, S10-S13), we identified a small set of robustly upregulated (179) and downregulated (260) genes across NC10-NC14 (Tables S16, S19, S21). Of the newly transcribed genes, we detected 111 genes with delayed transcription, including *frs* (Fig. 3F) and only

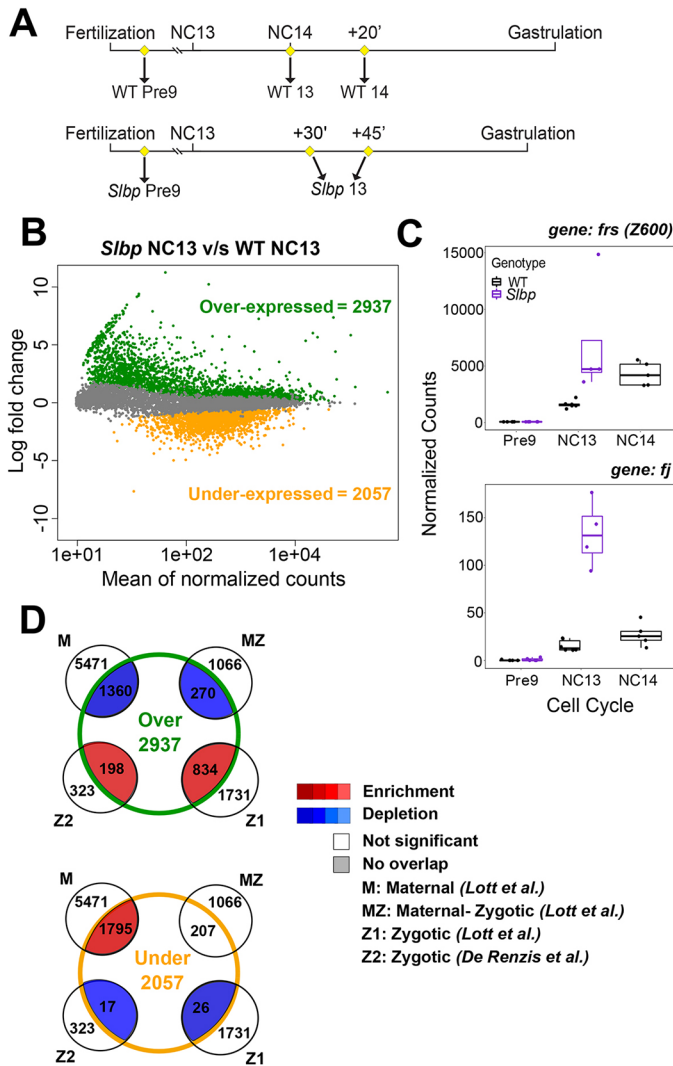


Fig. 2. Depletion of maternal histones results in an early ZGA.

(A) Schematic of embryo collection for *Slbp* RNA-seq. Pre-NC9 embryos were collected for both WT ($n=5$) and *Slbp* ($n=5$) to control for maternal loading. *Slbp* embryos were collected 30 or 45 min into NC13 ($n=4$) to ensure embryo health and phenotype and compared with NC-matched (mitosis of NC13) ($n=6$) or time-matched (20 min into NC14) ($n=5$) WT embryos. (B) When compared with WT NC13, 2937 genes are overexpressed and 2057 genes are underexpressed in *Slbp* embryos. (C) Traces from two overexpressed genes, *frs* and *ff*. Box plots: the box represents the interquartile range (ends of the box representing the upper and lower quartile), the central line is the median and the whiskers extend to the upper- and lowermost values. (D) Data from B compared with previous datasets. When compared with NC-matched controls, overexpressed transcripts are enriched for zygotically expressed genes and de-enriched maternal transcripts. Conversely, underexpressed genes are enriched for maternal transcripts and de-enriched for new zygotic transcripts. This pattern is consistent with premature ZGA. Comparisons with time-matched controls yield similar results (Fig. S3).

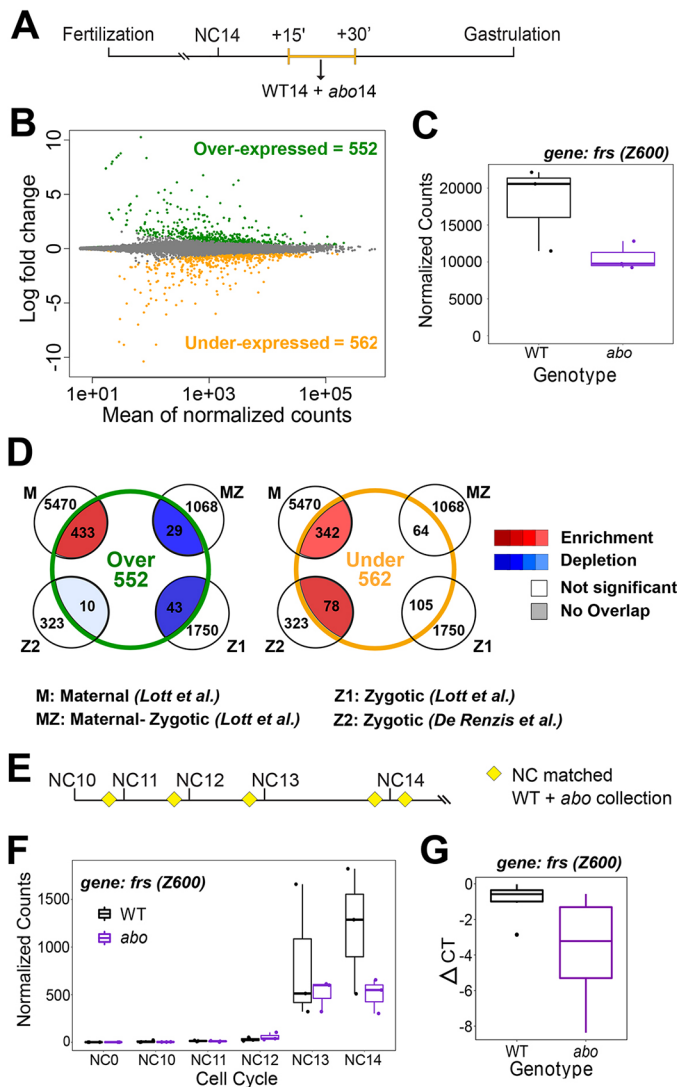


Fig. 3. Overexpression of maternal histones delays ZGA. (A) Schematic of embryo collection for B and C. Pooled *abo* ($n=3$) or WT ($n=3$) embryos were collected between 15 and 30 min into NC14. (B) 552 genes are overexpressed and 562 genes are underexpressed in *abo* embryos in NC14 compared with WT. (C) A representative trace of a zygotic gene (*frs*) that is differentially expressed in B. (D) Genes that are overexpressed in *abo* are enriched for maternal genes and those that are underexpressed are enriched for both maternal and zygotic genes. (E) Schematic of time-course embryo collection. Embryos were collected in the last 3 min of NC10-13 ($n=3$ per time point) and the first 3 min of NC14 ($n=3$). (F) A representative trace of a zygotic gene (*frs*) that is differentially expressed in NC14. (G) Single embryo qPCR of *frs* in NC14 *abo* and WT embryos (Δ CT normalized to *Rpl32*). Box plots: the box represents the interquartile range (ends of the box representing the upper and lower quartile), the central line is the median and the whiskers extend to the upper- and lowermost values.

37 that are upregulated (Table S14). We confirmed these results using qPCR (Fig. 3G). When compared with previous datasets, zygotic genes tend to be underexpressed, as was the case for the pooled *abo* dataset; however, the majority of these enrichments are not statistically significant (Fig. S6D). Nonetheless the majority of these underexpressed genes are expressed during NC14 in WT (Lott et al., 2011; Fig. S7). This geneset, in combination with the time-matched *Slbp* comparison, enables further examination of the chromatin features that underlie histone sensitivity for transcription independent of cell cycle changes.

Histone-sensitive genes contain specific chromatin features around the TSS

To identify chromatin features associated with histone sensitivity, we compared the presence of 143 modENCODE chromatin signals near the transcriptional start site (TSS \pm 500 bp) of genes whose expression is altered by changes in histone concentration independent of cell cycle time (Fig. S3, Figs 3E,4A). We found a clear pattern of unique chromatin features for the histone-sensitive genes, compared with all newly transcribed genes, that is highly similar between the histone over- and underexpression experiments (Fig. 4B, Tables S17,S18). The pioneer transcription factor ZLD, known to be important for nucleosome eviction during ZGA, is enriched in the promoters of histone-sensitive genes. Insulator proteins such as BEAF-32 and CP190 are depleted in histone-sensitive genes (Chen et al., 2013; Li et al., 2014). Promoters of histone-sensitive genes also show a strong reduction for SIN3A, a transcriptional repressor associated with cell cycle regulation (Pile et al., 2002; David et al., 2008). SIN3A is known to recruit HDACs to TSSs, and almost all HDACs also show significant de-enrichment at the TSSs of histone-sensitive genes (Kadosh and Struhl, 1998; Silverstein and Ekwall, 2005). Taken together, these marks make up a unique chromatin signature that may sensitize a locus to changes in histone concentration, as is likely for pioneer factors such as ZLD. Other aspects of this signature may indicate that these genes are subsequently subject to later developmental regulation, as indicated by H3K4me3 and H3K27me3 (Li et al., 2014; Chen et al., 2013).

DISCUSSION

In this study, we have demonstrated that histone concentration regulates the timing of the MBT in *Drosophila*, resulting in both early gastrulation and extra pre-MBT divisions from histone reduction and increase, respectively. Histone concentration also regulates ZGA. Thousands of genes are prematurely transcribed in histone-depleted embryos and hundreds of genes are delayed in histone-overexpressing embryos. The majority of these genes appear to be downstream of changes in cell cycle duration, suggesting a model in which histones directly regulate cell cycle progression. In other cell types, histone loss halts the cell cycle via accumulation of DNA damage and stalled replication forks (Ye et al., 2003; Prado and Aguilera, 2005; Groth et al., 2007; Gunjan and Verreault, 2003). In the early embryo, changes in histone availability may similarly create replication stress to directly or indirectly activate Chk1 as we have shown. In turn, Chk1 would inhibit Stg and/or Twine to slow the cell cycle (Di Talia et al., 2013; Farrell and O'Farrell, 2013; Deneke et al., 2016; Royou et al., 2008; Fasulo et al., 2012; Price et al., 2000; Ji et al., 2004; Stumpff et al., 2004; Shimuta et al., 2002). This mechanism is supported by previous observations that loss of zygotic histones causes the downregulation of Stg in the first post-MBT cell cycle (Günesdogan et al., 2014). In this case, the observed transcriptional changes would be independent or downstream of the altered cell cycle.

Alternatively, direct changes in transcription downstream of histone availability may feed into the cell cycle. In bulk, histone-sensitive transcripts might underlie the replication stress that has been previously proposed to slow the cell cycle at the MBT (Blythe and Wieschaus, 2015b). Consistent with this, the cell cycle lengthening and partial arrest phenotypes observed in mutant RNA Pol II embryos occur at a similar frequency to those we observe as the result of histone depletion (Sung et al., 2013). Another possibility is that specific histone-sensitive transcripts are

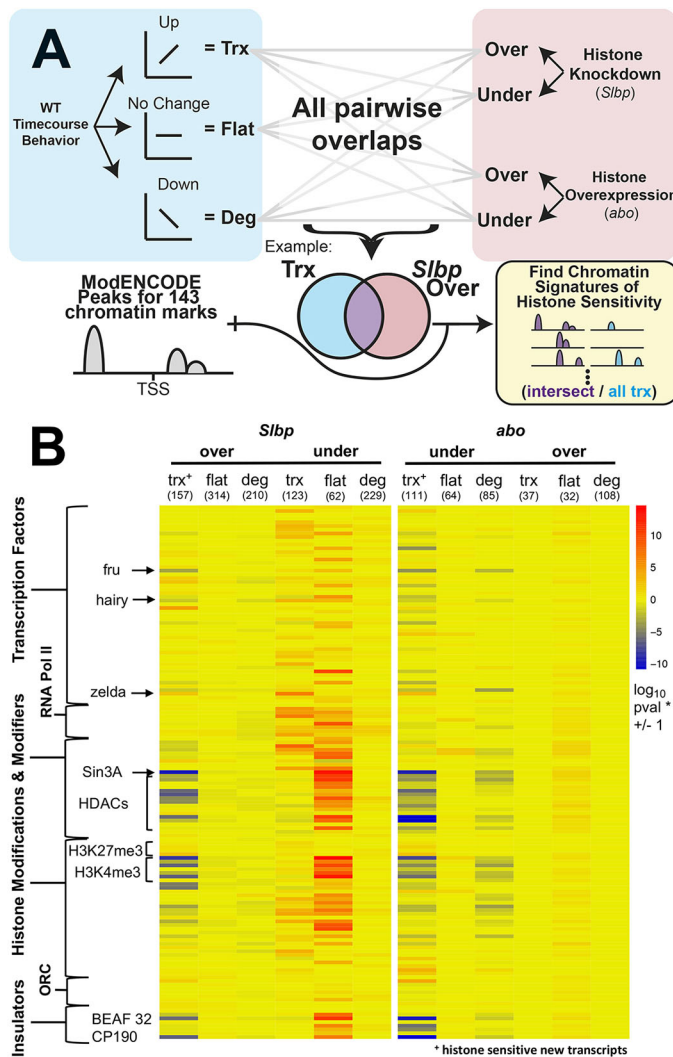


Fig. 4. Histone-sensitive transcripts share common chromatin features.

(A) Schematic of enrichment calculation. Genes were categorized as new transcription (Trx), flat or maternally degraded (Deg) based on their behavior between NC9 and NC14 in WT embryos (Fig. 2A). Next, significantly under- and overexpressed genes were identified from time-matched datasets for both *Slbp* (*Slbp* NC13 compared with WT NC14; Fig. S3D, Tables S2,S17) and *abo* (time-course; Fig. 3B, Tables S2,S18). These genes were then sorted based on WT behavior as Trx, Flat or Deg. This yielded 12 pairwise comparisons (e.g. new transcription and *Slbp* overexpressed, flat and *Slbp* overexpressed, etc.). Peak occupancy was calculated for all transcriptional start sites ± 500 bp in the genome from 143 modENCODE ChIP-seq and ChIP-chip datasets. Finally, enrichment for peaks from each modENCODE dataset was calculated using Fisher's exact test for each of the above 12 classes over the background class (e.g. the set of newly transcribed genes that were overexpressed in *Slbp* was compared with all newly transcribed genes). (B) Enrichments for 143 modENCODE datasets for six classes of genes, the expression of which was changed in *Slbp* or *abo* as described in A. Numbers below each heading denote the number of genes in each category. In both cases histone-sensitive new transcripts (the first column of each comparison, marked with +) were enriched for the pioneer transcription factor Zelda and H3K27me3 but de-enriched for class I insulator proteins (BEAF-32 and CP190), H3K4me3, *hairy*, *fruitless* (*fru*), SIN3A and its associated HDACs. ORC, origin recognition complex.

responsible for cell cycle elongation. One promising candidate for a histone-sensitive cell cycle regulator is the N/C ratio-sensitive CDK inhibitor *frs*, as zygotic transcription of *frs* plays a crucial role in stopping the cell cycle at the MBT (Großhans et al., 2003). In

contrast, *tribbles*, an N/C ratio-dependent inhibitor of Twine that has also been implicated in cell cycle slowing, does not show a consistent response between histone perturbations (Farrell and O'Farrell, 2013). In this previously proposed model, maternal histone stores may compete with pioneer transcription factors to set the timing of transcription initiation (Amodio et al., 2015; Joseph et al., 2017). Indeed, the central *Drosophila* pioneer transcription factor ZLD is enriched at the promoters of histone-sensitive genes. Moreover, we have identified a broader set of chromatin features that may sensitize individual loci to changes in histone concentrations. These include less obvious candidates for global early transcriptional regulators, such as SIN3A, HDACs and class I insulator proteins that may protect transcripts from changes in histone concentrations. Our work highlights the importance of histone concentration in regulating the timing of MBT and provides evidence that promoters of histone-sensitive genes possess a unique chromatin signature. However, future studies will be required to isolate the specific downstream effectors that respond to changes in histone concentrations in the early embryo.

MATERIALS AND METHODS

Drosophila stocks and genetic crosses

Slbp RNAi (BSC: 51171), *abo* (BSC: 2525), *OvoD/bTub85* (BSC: 2149) and *nls-RFP* (BSC: 31418) lines were obtained from the Bloomington *Drosophila* Stock Center. Oregon-R (Ore-R) and P(mat-tub-Gal4)mat67; P(mat-tub-Gal4)mat15 were a gift from Eric Wieschaus (Princeton University, USA). The Chk1 localization sensor was a gift from Stefano Di Talia (Duke University, USA). All fly stocks were maintained through standard methods at 25°C unless otherwise specified and grown on a standard cornmeal media. *Slbp* embryos were produced by crossing *nls-RFP*; P(mat-tub-Gal4)mat67; P(mat-tub-Gal4)mat15 driver virgin females to UAS-*Slbp* RNAi males at 18°C. The resulting female progeny were placed into cups with Ore-R males at 18°C and their embryos used for experiments. WT for *Slbp* experiments was *nls-RFP*; P(mat-tub-Gal4)mat67; P(mat-tub-Gal4)mat15 driver virgin females crossed to Ore-R males at 18°C. The resulting female progeny were placed into cups with Ore-R males at 18°C, and their embryos used as WT controls for *Slbp* experiments. *abo* embryos were produced by collecting *nls-RFP*; *abo*¹ homozygous females from an *nls-RFP*; *abo*¹/SM1 stock line. Females were placed into cups with Ore-R males at 25°C C, and their embryos were used for experiments. For *abo* control experiments *nls-RFP* females were collected and placed into cups with Ore-R males at 25°C C, and their embryos were used as WT. We found no significant deviations in cell cycle duration between WT embryos laid at 18°C or 25°C once imaging began at 22°C, indicating that any temperature-dependent effects on the cell cycle were mitigated by imaging at a constant temperature. Unfertilized embryos were collected from crossing *nls-RFP*; +; and *nls-RFP*; *abo*¹; homozygous virgin females to st1 β Tub85DD ss1 es/TM3, Sb1 males derived from the *OvoD/bTub85* (BSC: 2149). *Slbp* Chk1 sensor embryos were produced by crossing *nls-RFP*; Pmat-tub-Gal4)mat67; P(mat-tub-Gal4)mat15 driver virgin females to y,w; UAS-*Slbp* RNAi; Cdc25C[183-251]-EGFP males at 18°C. The resulting female progeny were placed into cups with Ore-R males at 18°C and their embryos used for experiments. WT for Chk1 sensor experiments was *nls-RFP*; P(mat-tub-Gal4)mat67; P(mat-tub-Gal4)mat15 driver virgin females crossed to y,w ; +; Cdc25C[183-251]-EGFP males at 18°C. The resulting female progeny were placed into cups with y,w males at 18°C, and their embryos used as WT controls for *Slbp* Chk1 experiments.

Microscopy

Embryos were dechorionated using 4% sodium hypochlorite and mounted on a 35 mm coverslip dish (MatTek) and covered with water. Cell cycle observations and RNA collections for *Slbp* and *abo* embryos were taken at 22°C using a Nikon Ti-E spinning disk confocal microscope with a 20 \times 1.3 NA air objective at 60 s/frame acquisition (1022 \times 1500 pixel area). WT and *Slbp* embryos expressing a Chk1 activity sensor were acquired at 24°C using

a Nikon A1R-Si HD confocal microscope with a 60×1.4 NA oil objective at 20 s/frame (500×248 pixel area). Images were processed with Nikon NIS-Elements and FIJI (Schindelin et al., 2012).

Cell cycle analysis

The duration of an NC was calculated from the number of frames between nuclear envelope breakdown in at least 50% of the nuclei in the embryo to 50% nuclear envelope breakdown in the next NC. Gastrulation was determined to be when multiple cells began movement away from the single tissue sheet.

Statistical significance for nuclear cycle duration between WT and *Slbp* embryos for each NC was determined using two-way ANOVA performed using R (3.4). Statistical parameters such as sample numbers, mean and adjusted *P*-value for multiple comparisons are included as follows: For NC12 – *Slbp* early ($n=7$, mean=23.9 min) versus WT ($n=6$, mean=16.7 min), $P=0.0000583$; *Slbp* early ($n=7$, mean=23.9 min) versus *Slbp* normal ($n=4$, mean=19.5 min), $P=0.0034035$; *Slbp* normal ($n=4$, mean=19.5 min) versus WT ($n=6$, mean=16.7 min), $P=0.1308121$. For NC13 – *Slbp* early ($n=4$, mean=86.8 min) versus WT ($n=7$, mean=25.6 min), $P<0.0000001$; *Slbp* normal ($n=4$, mean=33.3 min) versus WT ($n=7$, mean=25.6 min), $P=0.0056589$; *Slbp* early ($n=4$, mean=86.8 min) versus *Slbp* normal ($n=4$, mean=33.3 min), $P<0.0000001$.

Chk1 activity measurement

WT and *Slbp* embryos were imaged from NC11 to NC13 mitosis with a Chk1 activity sensor (Deneke et al., 2016). Chk1 activity was determined from the cytoplasmic-to-nuclear intensity ratio of the Chk1 localization sensor in a 150×75 pixel area. The nuclear signal was taken from segmented nuclei eroded by 2 pixels to ensure nuclear signal was analyzed. The cytoplasmic signal was taken from inverted nuclear masks dilated by 2 pixels. Four embryos per genotype were analyzed.

Single embryo qPCR

cDNA from RNA isolated from single embryos collected 3 min into NC14 was made with random primer mix using ProtoScript First Strand cDNA Synthesis Kit following the manufacturer's protocol (New England Biolabs, E6560L). qPCR was conducted on an ABI ViiA7 using the FG Taqman GEX master mix (Thermo Fisher Scientific, 4369016) and the following gene expression assays: *frs* (DM01822845; VIC) normalized to *RpL32* (DM02151887; FAM).

Western blotting

For western blotting, protein extracts were collected from WT, *Slbp* and *abo* embryos 55 min after pole bud formation, corresponding to early NC14 in the WT and individually stage-confirmed by halocarbon oil (Sigma-Aldrich, H8773). Embryos were washed with deionized (DI) water, dechorionated with 4% sodium hypochlorite, washed again with DI water, then lysed in ice-cold embryo lysis buffer (as per Günesdogan et al., 2014). Laemmli buffer (Bio-Rad Laboratories, 1610737) was added to each sample, and they were incubated at 95°C for 10 min. Five embryos were collected per sample. Proteins were separated on a TGX 12% acrylamide gel (Bio-Rad Laboratories), stain-free dye was crosslinked under UV for 1 min, and transferred to a low fluorescence PVDF membrane. Membranes were incubated overnight in rabbit anti-H3 antibody (1:2000; Abcam, ab1791) and mouse anti-H2B antibody (1:2000, Abcam, ab52484), washed, and then incubated for a minimum of 1 h in Alexa Fluor 647-conjugated goat anti-rabbit IgG antibody (1:2500; Invitrogen, A-21244) and/or Alexa Fluor 488-conjugated goat anti-mouse IgG antibody (1:2500; Invitrogen, A-11001). Fluorescence was detected using a gel imager (Bio-Rad ChemiDoc MP) and quantified in Image Lab. H2B and H3 signal were normalized to total protein using the bright ~45 kDa band in the stain-free channel which corresponds to vitellogenin. For the *Slbp* and WT comparison, the normalized H2B and H3 protein concentrations were averaged for each genotype and the average WT concentration for each protein was scaled to a value of 1. Error was calculated using a linear model in R (lm in base R) to account for gel differences and the mean genotype effects and the associated standard error were extracted. (Fig. S1A,B).

RNA collection – single embryo and pooled

Input RNA for RNA-seq and qPCR were collected from individual and pooled embryos laid by tightly staged WT (as defined above), *abo* homozygous and *Slbp* RNAi mothers gathered from apple juice agar plates with yeast. Individual embryos were placed into an RNase-free tube, were lysed and 100 µl lysis buffer (Applied Biosystems, KIT0214) was added. Embryos were then flash-frozen in liquid nitrogen and stored at –80°C. Pooled embryos were placed in 100 µl RNeasy Lysis Buffer (Qiagen, RNeasy Lysis Buffer) and stored at 4°C. When enough embryos were collected, RNeasy Lysis Buffer was removed and embryos were processed as above. See supplementary Materials and Methods for further details on RNA collection.

cDNA library preparation and sequencing

The integrity of total RNA samples was assessed on a Bioanalyzer 2100 using RNA 6000 Pico chips (Agilent Technologies).

Single WT and *Slbp* embryos

For single embryo WT and *Slbp* RNA samples, additional ribosomal RNA depletion was applied before the Smart-seq2 library preparation using the Ribo-Zero rRNA Removal (Human, Mouse, Rat) Kit (Illumina, MRZH11124). The cDNA samples, RNA-seq and libraries were examined on the Bioanalyzer (Agilent Technologies), using DNA HS chips for size distribution, and quantified by Qubit fluorometer (Invitrogen). Different DNA barcodes were added to each sequencing library, and the libraries to be sequenced together were pooled at equal molar amount. The RNA-seq libraries were sequenced on Illumina HiSeq 2500 Rapid flow-cells as single-end 75nt reads, following the standard Illumina protocol. Raw sequencing reads were filtered using Illumina HiSeq Control Software and only the Pass-Filter reads were used for further analysis.

Single, time-course WT and *abo* embryos

For RNA samples from single time-course WT and *abo* embryos, the poly-A-containing RNA transcripts were converted to cDNA using an oligo-dT adaptor and further amplified by PCR following the Smart-seq2 method (Picelli et al., 2014). Illumina sequencing libraries were made from the amplified cDNA samples using the Nextera DNA library prep kit (Illumina, FC-121-1031).

Pooled WT and *abo* embryos

For RNA samples from pooled WT and *abo* embryos, poly-A-containing RNA transcripts were enriched using oligo-dT bead and further converted to cDNA and Illumina sequencing library using PrepX RNA-seq library kit on the automated Apollo 324 NGS Library Prep System (Wafergen Biosystems) following the manufacturer's protocol.

Quantification and statistical analysis

Analyses on RNA-seq and modENCODE data were performed on the high-performance computing cluster (64-bit Springdale Linux) at the Lewis-Sigler Institute for Integrative Genomics, Princeton University, using the appropriate packages within the conda environment and package management system. All statistical analyses were performed using R (3.4), Bioconductor packages (3.8; Huber et al., 2015), and the conda package management system (4.5.11). In addition, we used custom Unix, Perl, Awk and Sed scripts as necessary. See supplementary Materials and Methods for greater detail on these analyses.

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Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualization: S.C., H.W., J.G., A.A.A.; Methodology: S.C., H.W., J.G., A.A.A.; Software: S.C.; Validation: H.W., J.G.; Formal analysis: S.C., H.W., A.A.A.; Investigation: H.W., J.G., A.A.A.; Data curation: S.C., H.W.; Writing - original draft: S.C., H.W., A.A.A.; Writing - review & editing: S.C., H.W., J.G., A.A.A.; Visualization: S.C., H.W., A.A.A.; Supervision: A.A.A.; Project administration: A.A.A.; Funding acquisition: A.A.A.

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Data availability

Raw sequences (FASTQ), Tab delimited raw Salmon quantification outputs, Tab delimited DESeq2 outputs and Tab delimited DESeq2 normalized gene counts for all genotypes and nuclear cycles have been deposited in GEO under accession number GSE136631.

Supplementary information

Supplementary information available online at <http://dev.biologists.org/lookup/doi/10.1242/dev.177402.supplemental>

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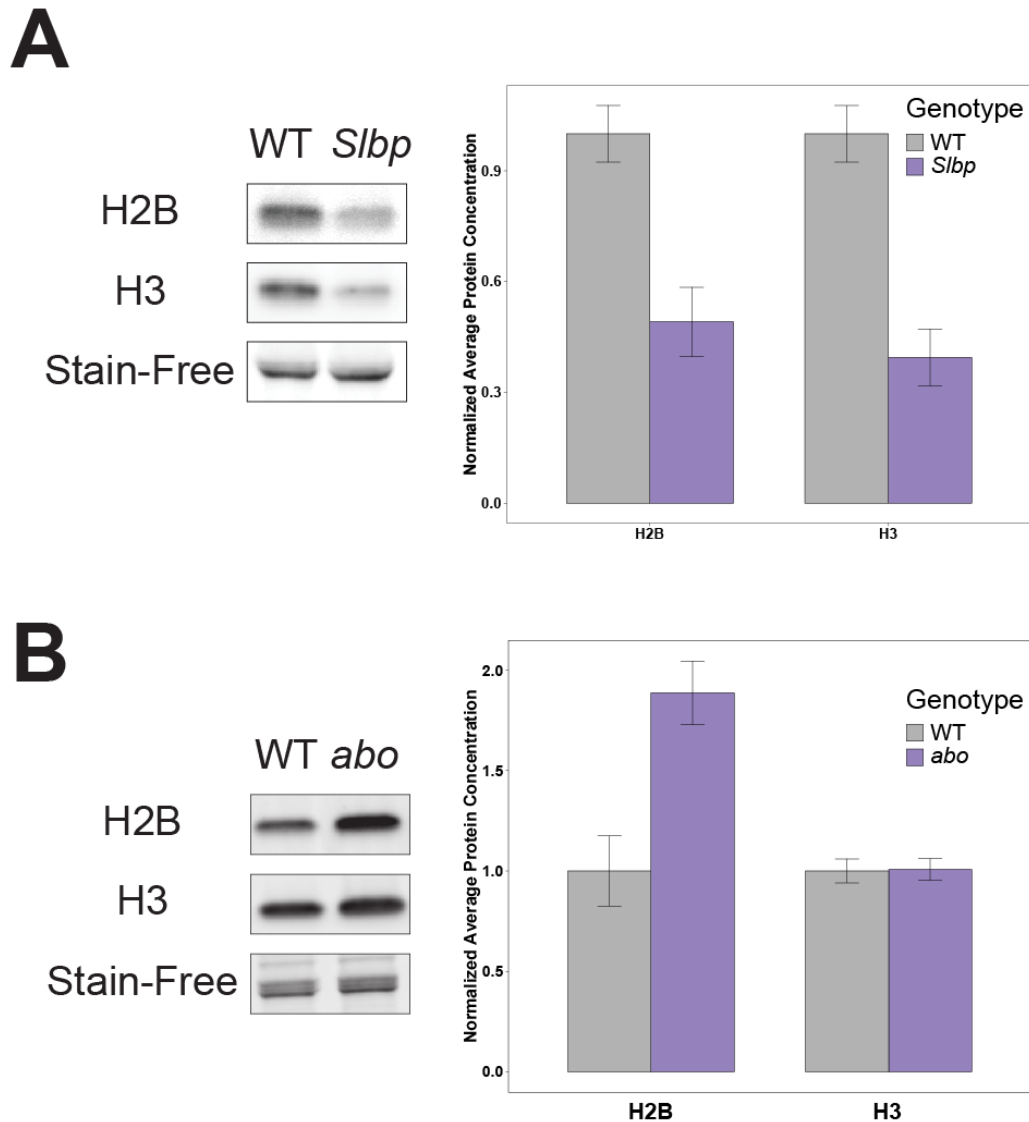
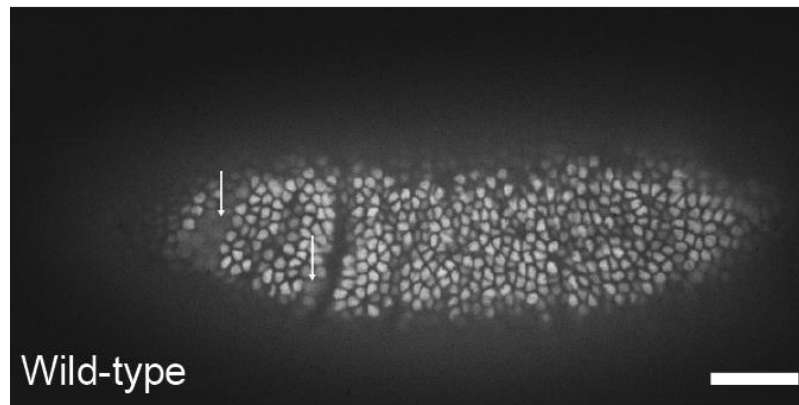


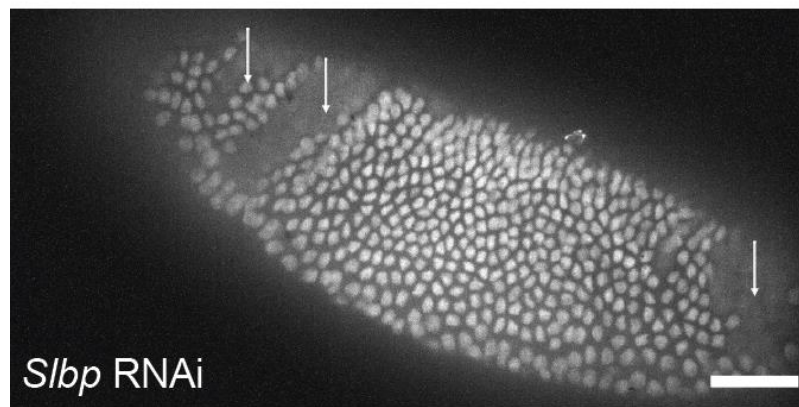
Figure S1. *Slbp* RNAi and *abo* embryos have altered histone protein levels

(A) Total H2B and H3 protein from WT and *Slbp* embryos collected 55 minutes after pole cell formation (early NC14 in WT) from western blotting with 5 embryos per lane. *Slbp* embryos have approximately a ~50% reduction in H2B protein and ~60% reduction in H3 protein (n=3). Bars represent mean and SEM. (B) Total H2B and H3 protein from *abo* embryos at collected 55 minutes after pole cell formation (early NC14 in WT) from western blotting. *abo* embryos have ~90% increase in H2B while total H3 was unchanged between WT (n=5) and *abo* (n=6) embryos. Bars represent mean and SEM. We note that the antibodies used for the assay cannot distinguish between the replication-coupled and replication-independent H3 variants and so the effect size on the replication-coupled H3 may be greater than reported here if there is compensation by the replication-independent variant.

A



B



C

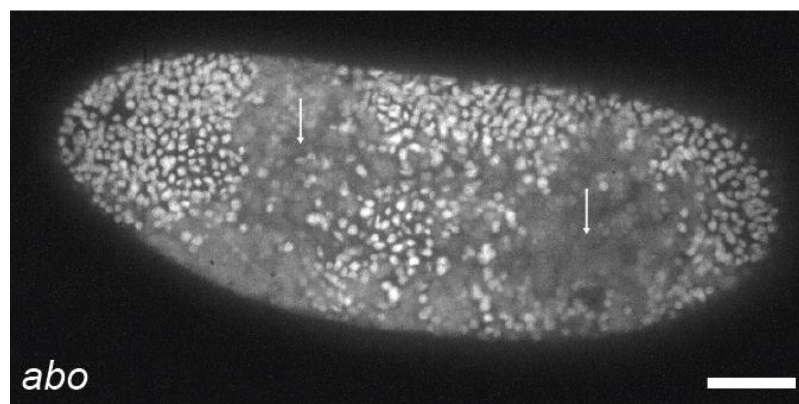


Figure S2. *S/bp* RNAi and *abo* embryos attempt post-MBT development

(A) WT embryo post-MBT after NC14. Mitotic domains and gastrulation movements form normally and with characteristic temporal and spatial dynamics. Arrows indicate mitotic domains. (B) *S/bp* embryo that enters gastrulation in NC13. Mitotic domain formation and gastrulation appear unaffected despite embryos having reduced numbers of cells. These embryos proceed through early-to-mid embryogenesis relatively normally but die before hatching. Arrows indicate mitotic domain formation. (C) *abo* embryos that enters gastrulation in NC15. *abo* embryos that undergo extra divisions still attempt post-MBT behaviors such as mitotic domain formation and gastrulation but are severely disrupted. Arrows indicate aberrant mitotic domain formation. Scale bar represents 50 μm .

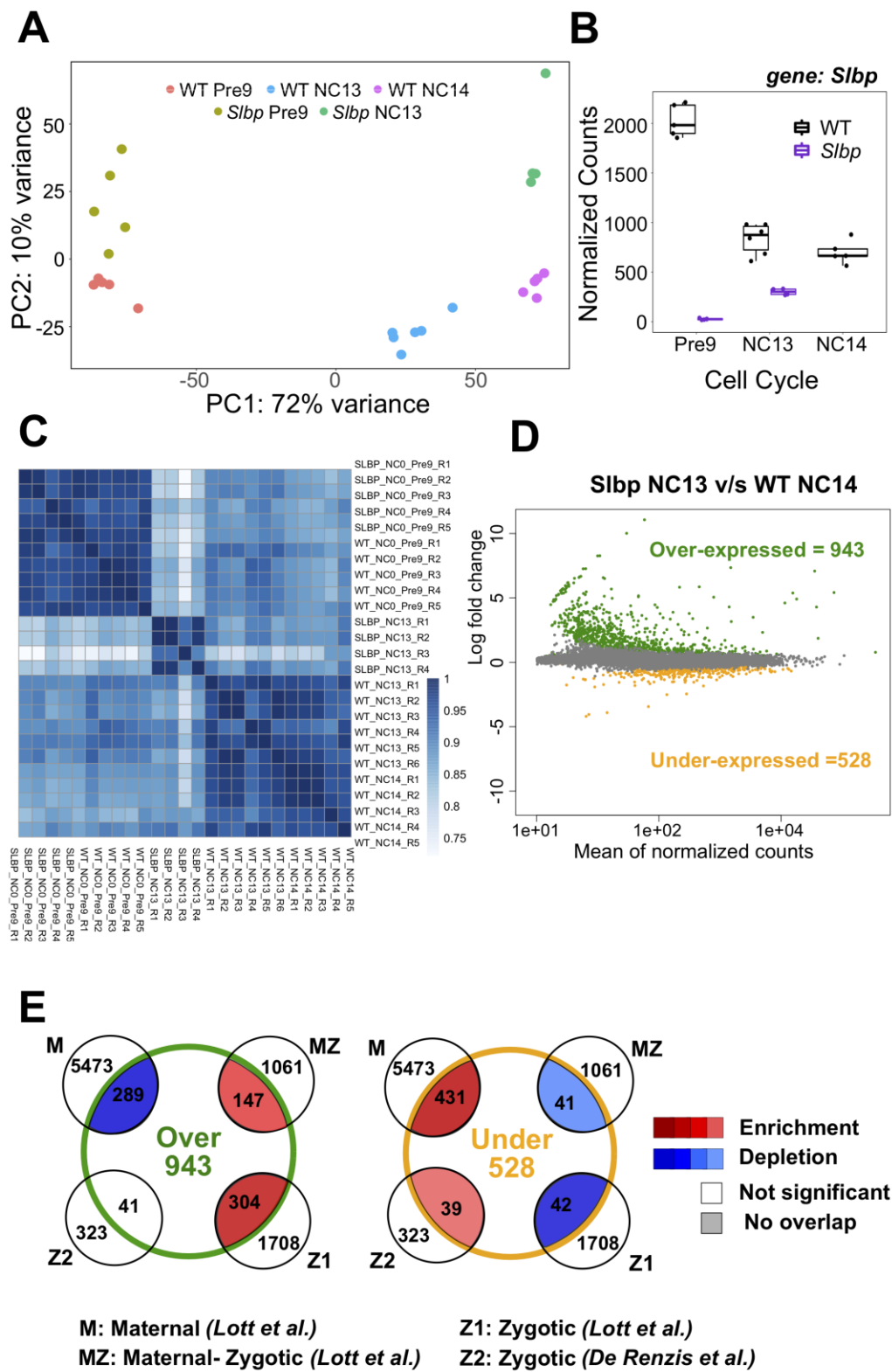


Figure S3. Depletion of maternal histones advance MZT

(A) Principal Components Analysis of *Slbp* and WT expression data demonstrating that global transcriptomic profile is shifted in *Slbp* NC13 as compared WT NC13 and is more similar to WT NC14 indicating that the onset of transcription is advanced in *Slbp* embryos. (B) Normalized counts of *Slbp* mRNA showing a reduction in *Slbp* mutants (n=5 in Pre9 and n=4 in NC13) as compared to WT (n=5 in Pre9, n=6 in NC13 and n=5 in NC14) (C) Spearman's correlation of expression data for WT and *Slbp* embryos at different timepoints. Replicates are more similar to each other than they are to other genotypes and timepoints. (D) When compared to time matched controls (NC14 +20') fewer genes are altered in expression. 943 genes are overexpressed and 528 genes are underexpressed, consistent with premature MZT in the *Slbp* embryos. (E) Data from C compared to previous datasets. Genes overexpressed compared to WT NC14 embryos show enrichment for zygotic transcripts while the underexpressed genes are enriched for maternal transcripts and show a slight enrichment for the De Renzis maternal genes.

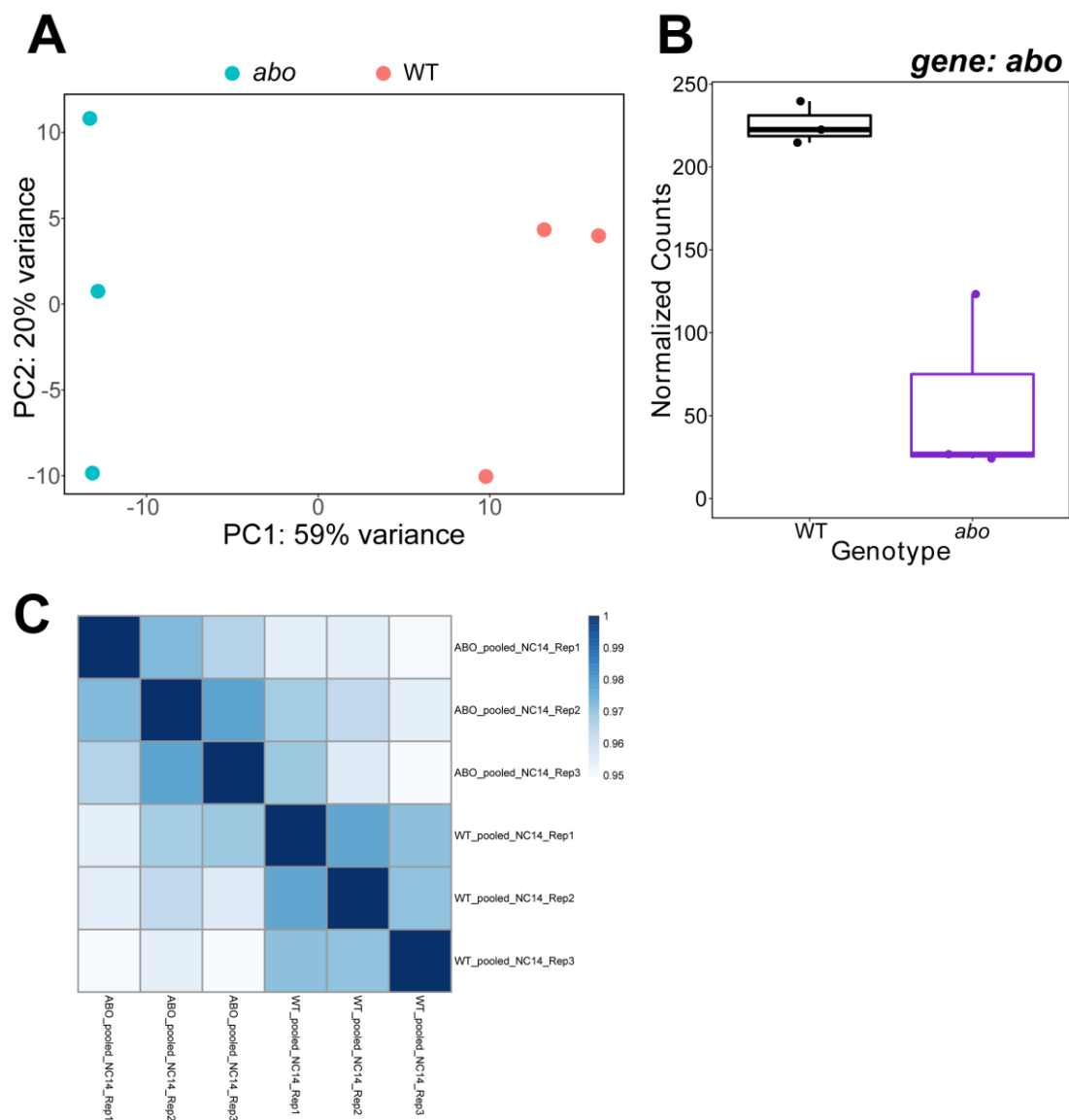


Figure S4. Global transcriptomic differences upon histone overexpression
(A) Principal Components Analysis between the pooled NC14 *abo* and WT expression data showing a clear difference in gene expression between genotypes at this developmental stage. (B) Normalized counts of *abo* mRNA showing reduction in *abo* mutants (n=3) as compared to WT (n=3). (C) Spearman's correlation of expression data for WT and *abo* embryos in NC14.

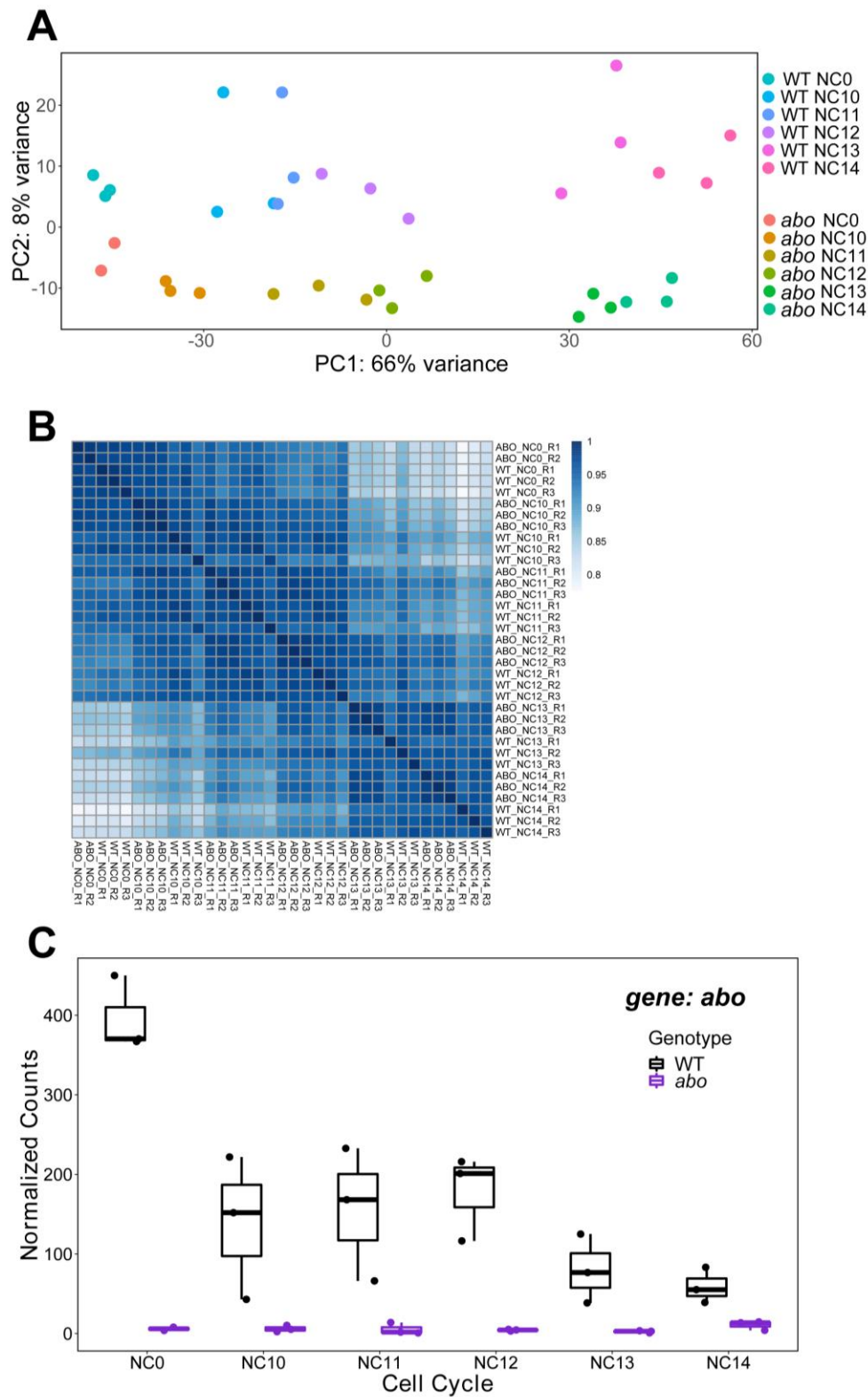


Figure S5. Timecourse transcriptomic differences upon histone overexpression

(A) Principal Components Analysis shows a clear separation across both cell cycle and genotype for the *abo* time-course experiment. (B) Spearman's correlation of expression data for WT and *abo* embryos across all timepoints. (C) Normalized counts of *abo* mRNA showing a reduction in *abo* mutants (n=2 at NC0 and n=3, all other time-points) as compared to WT (n=3, all time-points)

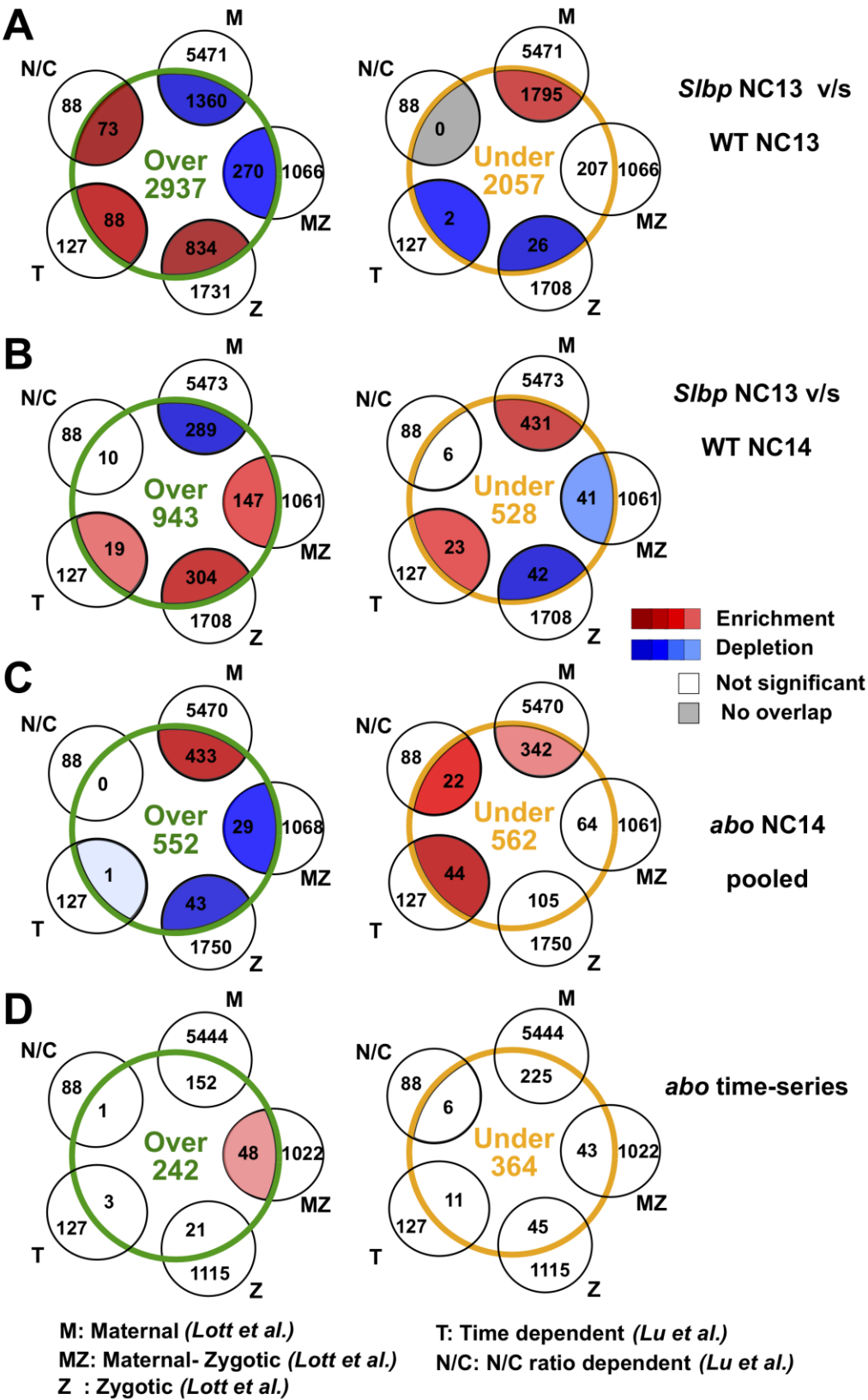


Figure S6. Histone sensitivity changes the onset of ZGA, but cannot explain previous N/C ratio dependent gene sets (A) *Slbp* NC13 compared to WT NC13. Data the same as Figure 2D except the De Renzis et al. (2007) maternal data has been split into N/C and time dependent genes as per Lu et al. (2009). Both N/C and time dependent genes are enriched in the overexpressed and depleted in the underexpressed categories, consistent with global advancement of the MZT. (B) *Slbp* NC13 compared to WT NC14. Data corresponds to Figure S3E. When controlled for time, enrichment of N/C ratio dependent genes is lost and time-dependent genes become enriched in both the over and under expressed gene sets. (C) *abo* NC14 compared to WT NC14. Data corresponds to Figure 3D. Both N/C and time dependent genes are enriched in the underexpressed category consistent with a global delay of ZGA. (D) *abo* time series comparison. Although not statistically significant, both N/C ratio and time dependent genes show an odds ratio >1 indicating these are also trending towards enrichment in the underexpressed gene set. Indeed, when the *abo* timecourse underexpressed genes are compared to De Renzis et. al (2007) zygotic genes dataset from which the N/C and time gene sets are derived the total zygotic overlap shows significant enrichment ($p=9.9e-4$) indicating that zygotic genes are better represented in the timecourse *abo* underexpressed group than the overexpressed set (Table S6).

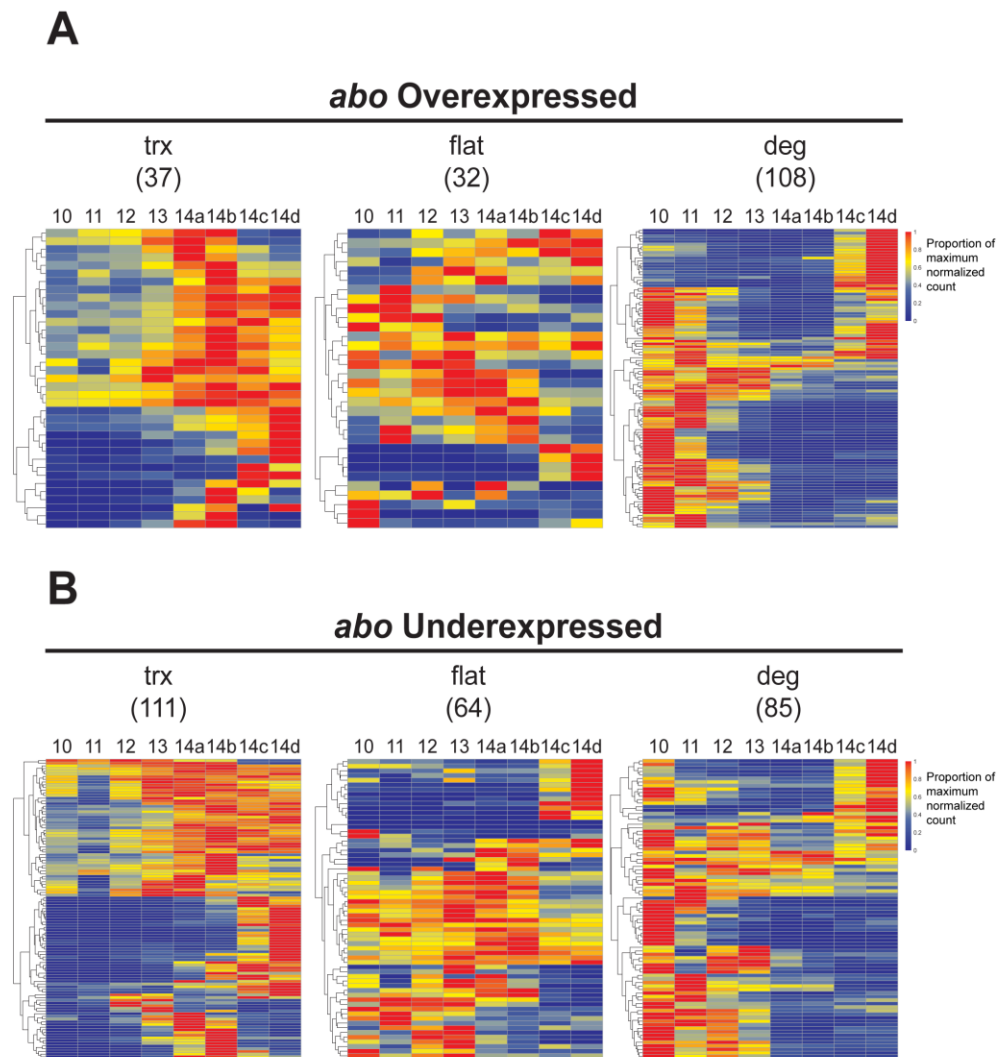


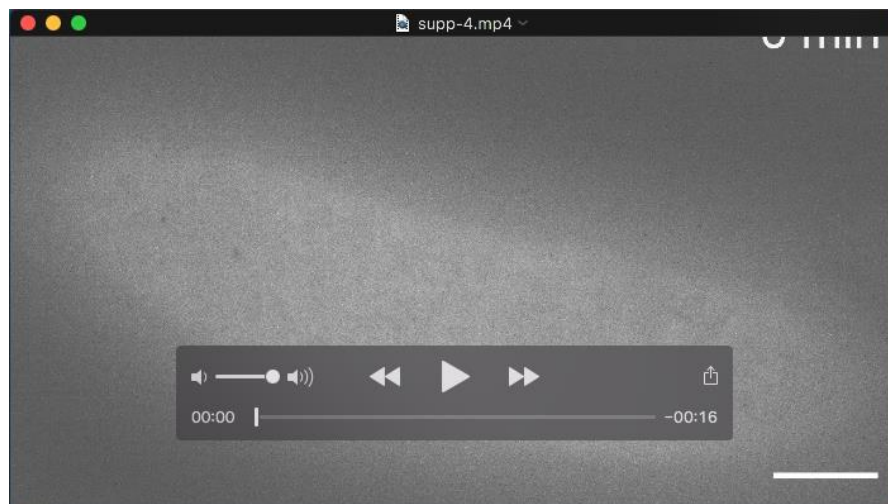
Figure S7. Genes that are knocked down by histone overexpression are expressed during WT MZT

(A) Overexpressed genes in the *abo* time series compared to longer time course from Lott et al. (2011). Each heatmap represents genes categorized as increasing (trx), remaining constant (flat), or decreasing (deg) from our control experiments (as in Figure 4). Numbers in parentheses indicate number of genes within the specified category. The majority of overexpressed genes are constant or downward trending, consistent with a delay in degradation of maternal products in histone overexpressing embryos. (B) *abo* underexpressed genes compared to Lott et al. (2011) Many of the underexpressed genes increase over time in WT supporting the idea there is a delay in zygotic transcription in *abo* embryos. Each of the heatmaps were generated by obtaining the median normalized count per gene per nuclear cycle and dividing these values by the maximum median normalized count value for that gene across all nuclear cycles. Thus, each of the cells in the heatmap is the proportion of the maximum normalized count for a given gene per nuclear cycle



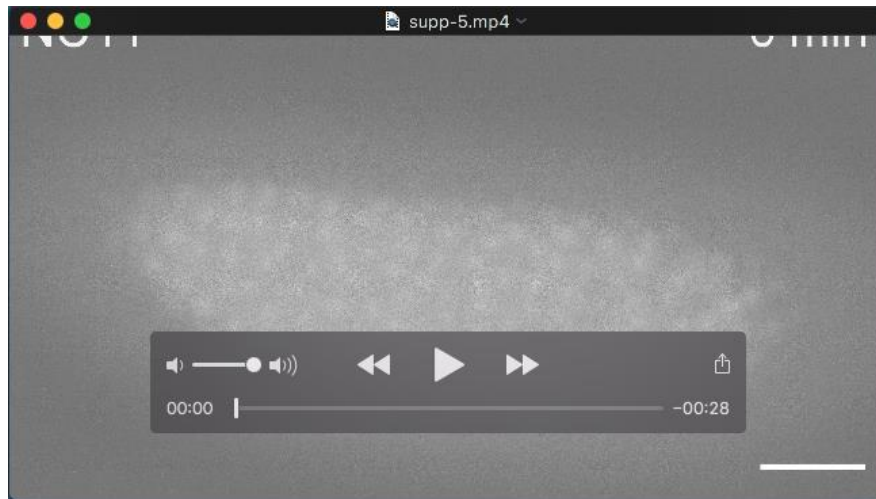
Movie 1. Representative *Slbp* embryo with full arrest in NC13

Slbp embryo has slower syncytial blastoderm nuclear cycles and ends the syncytial blastoderm cycles prematurely at NC13. It gastrulates one cell cycle early with about half the number of cells compared to WT. Still images from this movie are represented in Figure 1A. Scale bar represents 50 μm .



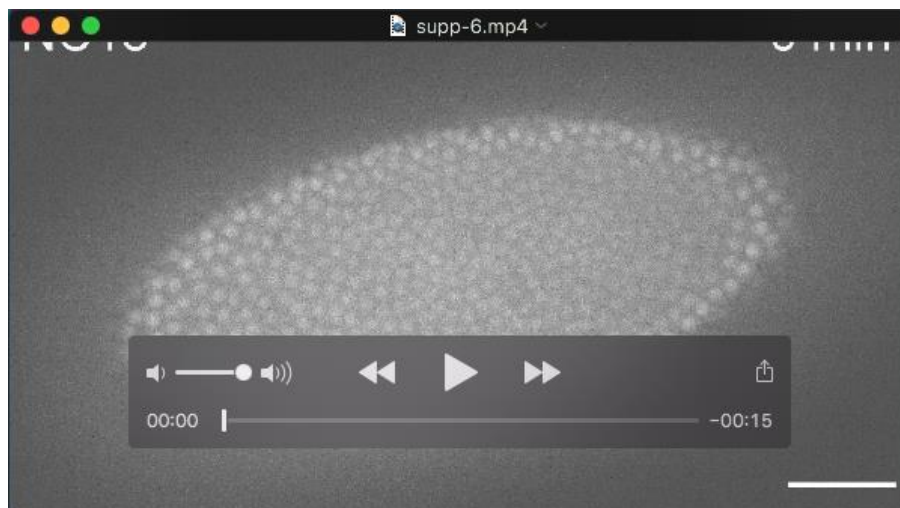
Movie 2. Representative *Slbp* embryo partially arrested in NC13

Slbp embryo has slower syncytial blastoderm nuclear cycles and partially arrests at NC13. It gastrulates normally with sections of the embryo in NC13 and NC14. Scale bar represents 50 μm .



Movie 3. Representative *abo* embryo with extra division

abo embryo has a shortened nuclear cycle 14 and goes on to have a 15th and 16th nuclear cycle before attempting gastrulation. Scale bar represents 50 μ m.



Movie 4. Representative *abo* embryo partially dividing into NC15

abo embryo has a normal NC14 before partially mitosing into NC15. It begins gastrulation while in the process of division and displays ectopic furrow formation. Scale bar represents 50 μ m.

Table S1. Syncytial blastoderm phenotypes of histone depletion and overexpression

Range of phenotypes during the syncytial blastoderm stage from *Slbp* and *abo* embryos. The percent of occurrence is noted in the table and total number in parenthesis beside this value. “No nuclei” is defined as no nuclei make it to the embryo’s surface during the course of live imaging which generally occurs ~90 mins after egg laying. “Catastrophic fallout” is defined as fallout which prohibits the embryo from successfully reaching the onset of gastrulation. “NC13” is defined as embryos which initiate their long cell cycle pause at NC13 instead of at NC14. “NC13/14” is defined as embryos which initiated their long cell cycle pause with part of the embryo in NC13 and another in NC14. “NC14” is defined as embryos which go through the syncytial blastoderm stage normally and initiate their cell cycle pause at NC14. “NC14/15” is defined as embryos which undergo a partial extra division before the onset of gastrulation. “NC15” is defined as embryos which undergo a complete extra division before the onset of gastrulation. We excluded from all analysis the ~70% of *Slbp* embryos which have defective blastoderm formation as have been previously reported in *Slbp* mutants and nulls (Sullivan et al., 2001; Iampietro et al., 2014). As previously described the majority of *abo* embryos arrest before the syncytial blastoderm stage (Tomkiel et al., 1991; Tomkiel et al, 1995). We therefore excluded these and *abo* embryos which undergo catastrophic fallout from analysis. However, of the ~40% of embryos that form blastoderm, ~6% do a complete extra division and another ~4% do a partial extra division.

[Click here to Download Table S1](#)

Table S2: Differentially expressed gene counts for all genotypes and cell cycles

The number of genes in each category for all differential expression analysis conducted in this manuscript.

[Click here to Download Table S2](#)

Table S3: Differentially expressed genes in all *Slbp* to WT comparisons

Table with results from the differential gene expression analysis (using DESeq2) including FDR adjusted p-values that were used to provide significance thresholds for all *Slbp* comparisons.

[Click here to Download Table S3](#)

Table S4: Gene lists for differentially expressed genes in all *Slbp* comparisons

Gene names and identifiers for all genes identified as differentially expressed in *Slbp* at either timepoint.

[Click here to Download Table S4](#)

Table S5: Normalized gene counts for *Slbp* and WT at each timepoint

Table with the normalized gene counts for *Slbp* and WT for each timepoint. These values are plotted in Figure 2B and S3D.

[Click here to Download Table S5](#)

Table S6: Comparison of previous datasets to differentially expressed genes identified in this study

Number of genes and overlaps in each category of comparison between all differentially expressed genes identified in this study and the Lott et. al (2011), De Renzis et. al. (2007), and Lu et. al (2009) datasets (gene lists for previous datasets included as Table S22). P-values generated by a two-sided Fisher's exact test. These numbers underlie the overlaps displayed in Figures 2D, 3D, S3E, and S6.

[Click here to Download Table S6](#)

Table S7: Differentially expressed genes in *abo* NC14 to WT NC14 comparisons

Table with results from the differential gene expression analysis (using DESeq2) including FDR adjusted p-values that were used to provide significance thresholds for the pooled *abo* comparison.

[Click here to Download Table S7](#)

Table S8: Gene lists for differentially expressed genes in *abo* NC14 to WT NC14 comparisons

Gene names and identifiers for all genes identified as differentially expressed for the pooled *abo* datasets.

[Click here to Download Table S8](#)

Table S9: Normalized gene counts for *abo* and WT at NC14

Table with the normalized gene counts for *abo* and WT for pooled dataset. These values are plotted in Figures 3B.

[Click here to Download Table S9](#)

Table S10: Differentially expressed genes in *abo* timecourse comparisons

Table with results from the differential gene expression analysis (using DESeq2) including FDR adjusted p-values that were utilized to provide significance thresholds for the *abo* timecourse comparison.

[Click here to Download Table S10](#)

Table S11: Gene lists for differentially expressed genes in *abo* timecourse comparisons
Gene names and identifiers for all genes identified as differentially expressed in the timecourse *abo* datasets.

[Click here to Download Table S11](#)

Table S12: Normalized gene counts for *abo* timecourse
Table with the normalized gene counts for *abo* and WT timecourse dataset. These values are plotted in Figure S5C.

[Click here to Download Table S12](#)

Table S13: Transcriptional trajectory of all genes detected between Pre9 to NC14 for the wild-type
Classification of all genes into “transcription”, “flat”, and “degraded” derived from Pre9 vs NC14 comparison in WT as described in figure 4A and Methods.

[Click here to Download Table S13](#)

Table S14: Differentially expressed gene distribution compared to WT transcriptional trajectory
Counts from advance gene-category classification representing histone manipulations v/s wild-type differential expression at a given stage compared to the temporal transcriptional profile in wild-type for the same set of genes (see Table 13 and Methods for more details). These numbers are used in Figures 4B and S7.

[Click here to Download Table S14](#)

Table S15: Differentially expressed genelist for genes in *Slbp* compared to WT transcriptional trajectory
Gene names and identifiers for all genes identified as differentially expressed in *Slbp* and classified into “transcription”, “flat”, and “degraded” categories derived from Pre9 vs NC14 comparison in WT. The comparisons to WT NC14 are used as the *Slbp* gene list in Figure 4B.

[Click here to Download Table S15](#)

Table S16: Differentially expressed genelist for *abo* timecourse compared to WT transcriptional trajectory
Gene names and identifiers for all genes identified as differentially expressed in *abo* timecourse and classified into “transcription”, “flat”, and “degraded” categories derived from Pre9 vs NC14 comparison in WT. These genes lists are used as the *abo* gene lists in Figures 4B and S7.

[Click here to Download Table S16](#)

Table S17: Comparison of modENCODE data and DE genes from *Slbp*

Table with gene numbers, overlaps and Fisher's exact test results for promoter peaks from different embryonic modENCODE ChIP-seq factors required for overlap with *Slbp* DE genes belonging to transcriptional categories defined as new transcription, flat, and degradation with appropriate background. These enrichments are plotted in Figure 4B.

[Click here to Download Table S17](#)

Table S18: Comparison of modENCODE data and DE genes from *abo* timecourse

Table with gene numbers, overlaps and Fisher's exact test results for promoter peaks from different embryonic modENCODE ChIP-seq factors required for overlap with *abo* DE genes belonging to transcriptional categories defined as new transcription, unchanged and degradation with appropriate background. These enrichments are plotted in Figure 4B.

[Click here to Download Table S18](#)

Table S19: Effects of thresholds cutoffs on result reproducibility

Table detailing the number of genes in each set for all comparisons with different combinations of p-value and expression fold change cutoffs. Note the marked loss of genes in the WT NC13 to WT NC14 comparison when expression cutoffs are applied.

[Click here to Download Table S19](#)

Table S20: Differentially expressed genes are also expressed in WT

Table showing the percent of genes in each comparison that were detected at all in the stage matched WT controls. The vast majority of genes in all comparisons are present in WT, just in different abundance indicating that our results are not the result of genome wide dysregulation.

[Click here to Download Table S20](#)

Table S21: Raw and mapped read counts

Table showing the number of raw reads and mapped reads for all datasets generated in this study.

[Click here to Download Table S21](#)

Table S22: Gene lists from literature datasets used in this study

Gene lists from Lott et. al (2011), De Renzis et. al (2007), and Lu et. al (2009) used for comparisons in this study. These gene lists underlie the overlaps displayed in Figures 2D, 3D, S3E, and S6.

[Click here to Download Table S22](#)

Supplementary Materials and Methods:

RNA collection - single embryo and pooled

Single WT and *Slbp* embryos:

Collections for RNA from single embryos were dechorionated with 4% sodium hypochlorite, washed with DI water, and then mounted in Nunc microwell trays with water (VWR, 470378). Nuclear stage was observed via RFP fluorescence with microscopy settings previously described. Nuclear stage was determined by tracking embryo development as nuclei first surface at nuclear cycle 10. Individual *Slbp* and WT embryos were confirmed to be in the desired nuclear cycle interphase or mitosis. Minimal fallout criteria were established to avoid massive downregulation of zygotic transition observed in *Slbp* mutant and null embryos due to embryonic death (Sullivan et al., 2001; Iampietro et al., 2014; Lefebvre et al., 2017). Embryos that displayed groups of nuclei falling out of the blastoderm surface during the nuclear divisions before collection were discarded. Individual embryos were placed into an RNase free tube, lysed with a sterile needle. Then 100 µl of lysis buffer (Applied Biosystems, KIT0214) was added, and embryos were flash-frozen in liquid nitrogen, and stored at -80°C. Six biological replicates were collected for WT 13 and WT 14. Five biological replicates were collected for *Slbp* NC13.

Individual Pre-NC9 WT and *Slbp* embryos were collected after one-hour laying, confirmed to be preblastoderm stage by visualization in halocarbon oil (Sigma, H8773), and processed as above. Five biological replicates were collected for each genotype.

Single, time course WT and *abo* embryos:

For individual *abo* and WT embryos, each embryo was mounted as described above and observed at the beginning of NC10. We discarded any embryo whose cell cycle times deviated from our standard times by more than 1 minute in any cycle before collection. These stringent conditions mean that we eliminated any *abo* embryo that would have undergone an extra division and that changes in cell cycle times cannot explain differences in gene expression. Individual embryos were placed into an RNase free tube, lysed with a sterile needle. Then 100 µl of lysis buffer (Applied Biosystems, KIT0214) was added, and embryos were flash-frozen in liquid nitrogen, then stored at -80°C. Individual unfertilized WT and *abo* embryos were collected after one-hour laying and processed as above. Three biological replicates were collected for each genotype at each time point.

Pooled WT and *abo* embryos:

Collections for RNA from pooled WT and *abo* embryos were placed under halocarbon oil to determine developmental stage. Embryos between 5 and 15 minutes into nuclear cycle 14 were washed with DI water, dechorionated with 4% sodium hypochlorite, and washed with DI water again. Dechorionated embryos were placed in 100 µl RNAlater (Invitrogen, AM7020) and stored at 4°C. Embryos continued to develop for an additional ~15 minutes in RNAlater that resulted in their final collection time of 15-30 minutes into NC14. When 43, 45, and 45 embryos were collected per genotype for each respective replicate RNAlater was removed and 100 µl of lysis buffer was added (Applied Biosystems, KIT02014). Embryos were lysed with a RNase free pestle, flash-frozen in liquid nitrogen, and stored at -80°C. RNA from all single and pooled embryos was isolated using PicoPure RNA Isolation Kit following the manufacturer's protocol (Applied Biosystems, KIT02014).

Quantification and Statistical Analysis

Data pre-processing

We split the barcodes using fastq-multx (1.3.1; Aronesty, 2013) and then utilized TrimGalore (0.5.0) using the default options to trim adapters. Trimmed reads were then quantified with Salmon (0.10.2; Patro et al., 2017) using the default options and the *Drosophila melanogaster* reference transcriptome (r6.19) obtained from flybase. The reference transcriptome was indexed using Salmon (0.10.2; Patro et al., 2017) with the default parameters prior to quantification (Table S21, S5, S9, S12).

Differential gene expression (DE) analysis

The counts generated by Salmon were utilized in DESeq2 (1.20.0; Love et al., 2014) to perform differential gene expression (DE) analysis comparing the “mutants” (*abo* or *Slbp*) to the wild-type per cell cycle. We used an FDR adjusted p-value cutoff ≤ 0.05 to determine significantly DE genes and log2fold-change greater than or less than 0 to assign genes to the significantly differentially over- or under-expressed categories respectively. In cases where multiple sequencing runs were used, the technical replicates (i.e. same library run on different flow-cells), were almost identical as we did not detect any DE genes between these (Fig S3A and C, Fig S4A and C, Fig S5A and B). Thus, we collapsed the technical replicates by summing the read counts prior to DE analysis. We additionally imposed a minimal threshold of detecting at least 1 read in 2 or more samples, after collapsing the technical replicates and prior to performing the DE analysis. To visualize similarities between biological replicates of all samples we used the plotPCA function in DESeq2 to visualize the samples in 2D spanned by their first two principal components (Figures S3A, S4A and S5A). We also plotted the Spearman’s rank correlation of the normalized counts as a heatmap to visualize correlations across all samples for a given experiment (Figures S3C, S4C and S5B). We did not detect a substantial difference in the total number of transcripts across different time points in a given experiment (Tables S5, S9, S12 and S21). However, due to degradation of maternal and accumulation of zygotic transcripts, not all transcripts are detected in all timepoints. To determine if the DE genes were maternal, zygotic, dependent on the N/C ratio or time dependent, we tested whether there was a significant overlap between the DE genes and relevant gene lists from literature via a two-sided Fisher’s exact test (Table S22). An overlap with a p-value ≤ 0.05 was considered as significant. To visualize the enrichment and de-enrichment of specific factors as a heatmap, we multiplied the log10 p-values by +1 for enrichment (shades of red) and -1 for de-enrichment for plotting (shades of blue) based on the magnitude of the odds ratio (>1 or <1 respectively) in Figures 2D, 3D, S3E and S5. See table S6 for all comparisons.

Advance gene-category classification

To generate a robust set of differentially expressed genes across multiple cell cycles we utilized Fisher’s p-value combination method implemented in the metaRNASeq R package (1.0.2; Rau et al., 2014), to combine the p-values from all of the independent tests between the mutant and wild-type per cell cycle and generated a global FDR adjusted p-value for each gene across all cell cycles. We then assigned a gene significantly DE if it had a global FDR adjusted p-value ≤ 0.05 . Subsequently, we inferred the signs of the p-value combined DE genes by comparing with differentially up or down-regulated genes identified at each cell cycle (Table S3, S7, and S10). We excluded any gene that was identified as significantly DE in the global analysis but were not identified as DE in any of the individual cell cycle. We further excluded all genes DE in the unfertilized embryo comparison from the larger set of DE genes, since these would reflect differences in maternal loading. To further classify whether DE genes were DE in mutants due

to excess transcription or degradation as compared to wild-type across time, we identified the genes that were upregulated, down-regulated or unchanged (flat) between NC14 and unfertilized embryo in only the wild-type, based on an FDR adjusted p-value ≤ 0.05 (Table S13). Any gene that is up-regulated in this window, we classify as “new transcription” and genes down-regulated as “degradation”. These two categories largely meet expectations for gene membership. For instance, we find previously identified zygotic genes (De Renzis et al., 2007; Lott et al., 2011) over represented in the new transcription category. However, our classification is more permissive as transcripts that increase over time are included in the new transcription category regardless of the maternal contribution. We then determined how many of the DE genes (up/ down in mutant v/s wild-type) overlapped with the “new transcription” (trx), “degradation” (deg) and “flat” category of genes across time, as inferred from the wild-type data (Table S14, S15, and S16). Thus, genes that are differentially down-regulated in the mutant v/s wild-type comparison but overlap with “new transcription” category are those that have reduced transcription in the mutant over time as compared to the wild-type. While genes that are differentially up-regulated in the mutant v/s wild-type comparison but overlap with “degradation” category are those with increased stability in the mutant over time relative to the wild-type.

modENCODE comparisons

Chromatin signature identification using modENCODE ChIP-- data enrichment at promoters To understand if genes belonging to the various DE gene-categories were enriched for other chromatin features, we performed an enrichment analysis using all modENCODE ChIP-seq and ChIP-chip datasets at developmental stages including 0-4h, 0-8h, 0-12h and 0-24h post-fertilization (modENCODE Consortium et al., 2009). While the 0-4h developmental stage provides the most appropriate comparison for our experiment, we wanted to make sure that our analysis captured all of the early embryogenesis related chromatin signatures. We converted the ChIP-peak coordinates to dm6 using a perl script provided by flybase for coordinate conversion and further re-annotated these peaks using ChIP-seeker R package (1.16.1; Yu et al., 2015), with TSS +/-500bp as being promoters. We subsequently extracted all the genes for which there is any ChIP-peak within the promoter region and performed a gene-set enrichment analysis with genes we identified from the relevant RNA-seq analysis. Enrichment analysis was performed via a two-sided Fisher’s exact test in R with the appropriate gene categories. For example, we compared whether the genes that are underexpressed in a given histone manipulation v/s wild-type but are increasing over time in WT (i.e., *Slbp* or *abo* Under expressed and “new transcription” category as defined above) were enriched for the presence or absence of promoter peaks as compared to a background of all newly transcribed genes. An overlap with an FDR adjusted p-value ≤ 0.05 was considered as significant. To visualize the enrichment and de-enrichment of specific factors as a heatmap, we multiplied the log10 p-values by +1 for enrichment (shades of red) and -1 for de-enrichment for plotting (shades of blue) based on the magnitude of the odds ratio (>1 or <1 respectively) in Figure 4B. See supplemental tables S17 for *Slbp* and S18 for *abo* comparisons.

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