Evolutionarily Conserved Roles for Apontic in Induction and Subsequent Decline of Cyclin E Expression

HIGHLIGHTS

- Mutual activation of apt and e2f1 promotes rapid induction of CycE at S phase entry.
- Apt also up-regulates Rbf1, but Rbf1 is inactivated through phosphorylation by Cdk2.
- After initiation of S phase, Rbf1 becomes active and represses cycE.
- Apt governs both induction and subsequent repression of cycE.

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Evolutionarily Conserved Roles for Apontic in Induction and Subsequent Decline of Cyclin E Expression

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SUMMARY
Cyclin E is a key factor for S phase entry, and deregulation of Cyclin E results in developmental defects and tumors. Therefore, proper cycling of Cyclin E is crucial for normal growth. Here we found that transcription factors Apontic (Apt) and E2f1 cooperate to induce cyclin E in Drosophila. Functional binding motifs of Apt and E2f1 are clustered in the first intron of Drosophila cyclin E and directly contribute to the cyclin E transcription. Knockout of apt and e2f1 together abolished Cyclin E expression. Furthermore, Apt up-regulates Retinoblastoma family protein 1 (Rbf1) for proper chromatin compaction, which is known to repress cyclin E. Notably, Apt-dependent up-regulation of Cyclin E and Rbf1 is evolutionarily conserved in mammalian cells. Our findings reveal a unique mechanism underlying the induction and subsequent decline of Cyclin E expression.

INTRODUCTION
As a key factor for S phase entry, Cyclin E (CycE) is crucial for both mitotic and endocycling cells (Dulic et al., 1992; Knoblich et al., 1994; Lilly and Spradling, 1996). Previous studies have shown that transcription factor E2f, a heterodimer of E2f1 and Dp, plays an important role in cycE expression in normal cell cycle and endo-cycle (De Veylder et al., 2002; Duronio et al., 1995; van den Heuvel and Dyson, 2008; Zielke et al., 2011). However, Drosophila e2f1 null mutant can survive to the third-instar larval stage and residual S phase occurs in the mutant cells (Duronio et al., 1998; Royzman et al., 1997), suggesting an involvement of another transcription factor in S phase entry.

One candidate for such factor is Apt. Drosophila Apt (also termed Trachea defective, Tdf) is a DNA-binding transcription factor that is involved in the development of multiple organs and tissues, such as tracheae, head, heart, ovary, stem cell, nervous system, and imaginal discs (Eulenberg and Schuh, 1997; Gelion et al., 1997; Lie and Macdonald, 1999; Liu et al., 2003, 2014; Monahan and Starz-Gaiano, 2016; Shen et al., 2018; Starz-Gaiano et al., 2008; Su et al., 1999; Wang et al., 2017). We have found that Apt directly regulates the expression of cycE during the development of imaginal discs (Liu et al., 2014; Wang et al., 2017). Therefore, Apt might participate in the expression of cycE in other tissues also. Besides, Apt can suppress tumor metastasis, and the human homolog of Apt, FSBP, is a cancer-related factor (Lau et al., 2010; Woodhouse et al., 2003).

Although CycE is crucial for S phase entry, it should decrease subsequently for progression of the cell cycle. Rbf1 is a key player in the decline of CycE expression (Cayirlioglu et al., 2003; Korenjak et al., 2012; van den Heuvel and Dyson, 2008; Weng et al., 2003). During S phase entry, Rbf1 is inactivated by phosphorylation with Cyclin-dependent kinase 2 (Cdk2) but becomes active by de-phosphorylation after initiation of S phase (Du et al., 1996; Edgar and Orr-Weaver, 2001; van den Heuvel and Dyson, 2008). The activated Rbf1 binds to E2f1 and also forms another complex containing E2f2 and Dp to repress E2f1-target genes including cycE and many other genes (Cayirlioglu et al., 2003; Korenjak et al., 2012; van den Heuvel and Dyson, 2008; Weng et al., 2003). Retinoblastoma protein (Rb), a mammalian counterpart of Rbf1, promotes chromatin compaction for transcriptional silencing by interaction with chromatin regulators such as histone deacetylases and histone methyltransferases (Brehm et al., 1998; Nielsen et al., 2001; Talluri and Dick, 2012). Therefore, Rbf1 is also expected to participate in chromatin compaction for silencing.
Here we provide evidence that both Drosophila Apt and mouse FSBP play important roles in the induction of CycE and up-regulation of Rbf1 for proper chromatin compaction. Mechanistically, we showed that Apt and E2f1 mutually activate the expression of each other to induce cycE for S phase entry in the salivary gland. Furthermore, we observed that the binding motifs of Apt and E2f are clustered in the first intron of cycE. Based on the results of chromatin immunoprecipitation (ChIP) and transgenic reporter assays, we found direct contribution of the Apt-binding sites and the E2f1-binding sites to the transcription in the salivary gland. Moreover, we also found that Apt up-regulates Rbf1 to direct proper chromatin compaction for transcriptional silencing. Finally, we demonstrated evolutionary conservation of these mechanisms in mammalian cells.

RESULTS

Apt and E2f1 Activate Expression of Each Other

To investigate the function of Apt in endoreplication of the salivary gland, we first compared the expression of Apt and E2f1 proteins by immunostaining (Figure 1A). Endocycle in the salivary gland proceeds asynchronously, and hence each cell resides in various phases of endocycle. E2f1 peaks at S phase entry and declines after initiation of S phase (Zielke et al., 2011). According to the oscillation of E2f1 during endocycle, some cells expressed E2f1 strongly, whereas other cells expressed weakly. Intriguingly, we noticed that the expression of Apt exhibits a similar pattern as that of E2f1. The observed tight correlation between the levels of Apt and E2f1 proteins suggests almost-synchronous oscillation of Apt and E2f1 during endocycle.

As Apt and E2f1 are transcription factors, the strong correlation between the levels of Apt and E2f1 could be due to the interdependence of the apt and e2f1 expression. To test the possibility, we analyzed mRNA levels of apt and e2f1 in the salivary gland by RT-qPCR. To compare the mRNA levels among samples with different genome dosages, each mRNA level was normalized to that of b-tubulin mRNA. RNAi knockdown of e2f1 using a dpp-GAL4 driver (Figure S1A) decreased the expression of apt, and vice versa (Figure 1B). Furthermore, the expression of Apt and E2f1 proteins were dependent on each other (Figure S1B). These data demonstrate mutual activation of apt and e2f1. The positive feedback between apt and e2f1 would support their rapid and robust transaction. To examine whether these activations are direct or not, we searched for E2f1- and Apt-binding motifs in the apt or e2f1 promoter. E2f1-binding sites were found in the apt promoter region, suggesting that E2f1 might directly activate apt transcription. We used the 1.5-kb promoter region containing the E2f1-binding sites (Figure S2) to verify this possibility through transgenic reporter assays. As we expected, wild-type reporter gene was expressed in the salivary gland (Figure 1C), whereas the expression level of the reporter gene significantly decreased in the E2f1-binding site mutant line (Figures 1C and 1D). These results demonstrate that E2f1 can directly activate apt transcription in the salivary gland. Because the Apt-binding site was not found in the e2f1 promoter region, the apt-mediated activation of e2f1 might be indirect.

Both Apt and E2f1 Are Required for CycE Expression and Endoreplication

Considering the tight correlation between the expression of Apt and E2f1, it is most likely that Apt is involved in endocycle together with E2f1. To test the possibility, we induced apt null mutant clones in embryonic salivary glands where cells still undergo mitosis and observed the glands at the third-instar larval stage after many rounds of endoreplication in control cells. Compared with control cells, apt-knockout cells showed obvious decrease in DAPI fluorescence (Figures 2A and 2B). The decreased DAPI fluorescence indicates the role for Apt in endoreplication. In addition to the decreased DAPI staining, loose chromatin appearance was observed in apt-mutant clone cells (Figure 2A). We will refer to the phenotype in the later section.

In agreement with the previous studies (De Veylder et al., 2002; Duronio et al., 1995; van den Heuvel and Dyson, 2008; Zielke et al., 2011), knockdown of e2f1 decreased the expression of cycE in the salivary gland (Figure 1B). Furthermore, knockdown of apt also reduced the expression of cycE. Having established that both Apt and E2f1 are cycE activators, we compared the DNA content and CycE protein level among control cells, apt-knockout cells, e2f1-knockout cells, and apt- and e2f1-double knockout cells. The DNA content decreased clearly in apt- or e2f1-knockout clone cells (Figures 2A, 2B, 2D and 2H). Double knockout of apt and e2f1 completely blocked endoreplication (Figures 2E, 2E’ and 2H). Knockout of either apt or e2f1 significantly decreased the CycE expression, but residual CycE protein was still detectable (Figures 2C and 2F). Upon double knockout of apt and e2f1, the expression level of CycE reduced below the...
These data collectively demonstrate that both Apt and E2f1 are required for proper CycE expression and endoreplication in the salivary gland.

**Apt and E2f1 Can Directly Activate cycE Transcription**

As Apt and E2f1 activate the expression of each other and both Apt and E2f1 are required for the CycE expression, effect of e2f1 null mutation on the CycE expression is a combination of a direct effect due to the absence of E2f1 and an indirect effect due to the reduced Apt level. To address the direct contribution of Apt or E2f1 to cycE transcription, we focused on cis-regulatory elements of cycE. Expression of cycE is regulated by complex tissue-specific cis-elements (Jones et al., 2000). Although cis-elements for the expression in the salivary gland have not been reported, we found a clustering of two adjacent E2f1-binding motifs and four Apt-binding motifs in the first intron of cycE (Figure 3A). To test whether Apt or E2f1
Figure 2. Apt Is Required for Endoreplication and CycE Expression in the Salivary Gland

(A) An apt-mutant clone (marked by white dotted lines) was stained with DAPI. n = 12 with all samples showing the represented phenotype. Scale bar, 20 μm.

(B) Quantification of DAPI fluorescence from apt-mutant clone cells and neighboring control cells. Data were presented as mean ± SD. n = 4 cells for apt-mutant clones and 10 cells for control. ***p < 0.001 versus the control (Student’s t test).

(C) Immunofluorescence staining with anti-CycE antibody and DAPI in apt-mutant clones. Arrows show reduced expression of CycE in apt-mutant clones. n = 11 with all samples showing the represented phenotype. Scale bar, 20 μm.

(D) e2f1-knockout cells (marked by red arrowheads) showed lower ploidy. n = 7 with all samples showing the represented phenotype. Scale bar, 20 μm.

(E) An apt- and e2f1-double knockout cell (marked by yellow arrowheads) showed almost no endoreplication. n = 4 with all samples showing the represented phenotype. Scale bar, 20 μm.

(F) Close-up image around the yellow arrowhead in (E). Scale bar, 20 μm.

(G) CycE was decreased but still detectable in an e2f1-mutant cell (marked by red arrowheads). n = 4 with all samples showing the represented phenotype. Scale bar, 20 μm.

(H) apt- and e2f1-double mutant cell (marked by yellow arrowheads) abolished CycE expression. n = 3 with all samples showing the represented phenotype. Scale bar, 20 μm.
Apt Up-regulates Rbf1 and Directs Proper Chromatin Compaction

e2f1 mutant cells induced in the salivary gland displayed small nuclei with low DNA content (Figure 2D). This is exactly expected from the reduced endoreplication. By contrast, nuclei of apt mutant cells were abnormal. Despite the reduced level of endoreplication, the size of nuclei in the apt mutant clone cells was comparable to that of control cells (Figure 2A). The ratio of nuclear size to DNA content was ~6.5 times higher in apt-mutant cells than that in control cells (Figure 4A). A higher-magnification image of apt mutant nuclei exhibited intra-chromosomal chromatin de-compaction and large inter-chromosomal spaces compared with control nuclei (Figure S5). As the loose chromatin is associated with increased transcription activity, loss of apt function would induce de-repression of multiple genes. Indeed, apt-knockout cells exhibited many signals of transcribing RNA polymerase II, under the conditions where the signals were barely detectable in control cells (Figure 4B). Then, what is a global repressor that governs the silencing of many genes in the downstream of Apt? One candidate is Rbf1, because it forms a complex with E2f1 and another complex including E2f2 and Dp to repress E2f-target genes and many other genes (Cayirlioglu et al., 2003; Korenjak et al., 2012; van den Heuvel and Dyson, 2008; Weng et al., 2003). Consistent with this idea, we observed large nuclei with de-compact ed chromatin upon RNAi knockdown of rbf1 (Figure 4C). Therefore, it is possible that Apt up-regulates rbf1, and hence the apt mutant cells exhibit large nuclei with de-compact ed chromatin. To test the possibility, we analyzed the expression of rbf1 in the salivary gland by RT-qPCR. As shown in Figure 4D, we observed a significant reduction in the expression of rbf1 and e2f2 upon RNAi knockdown of apt leaving the dp expression unchanged. We also examined the expression of Rbf1 protein in apt-knockout or apt-overexpressing cells. Compared with control cells, apt-mutant clone cells showed decreased expression of Rbf1 protein, whereas apt-overexpressing cells exhibited highly compact chromatin with increased Rbf1 protein levels (Figures 4E and 4F). Consistently, we observed de-repression of Rbf1-target genes, such as CG4679, gigas, diap3, and lpp upon knockdown of apt (Figure 4D). Importantly, overexpression of Rbf1 suppressed chromatin de-compaction upon RNAi knockdown of apt (Figure 4G). Based on these results, we reasoned that the nuclear defects in the apt-knockout cells are due to release from Rbf1-mediated chromatin compaction and de-repression of many Rbf1-target genes.

There exists a single Apt-binding motif at 156 nucleotides upstream of the transcription start site of rbf1. ChIP assays showed occupancy of Apt on the motif (Figure 4H). These data suggest that rbf1 is a direct target of Apt.

Figure 2. Continued

(H) Quantification of DAPI fluorescence in e2f1-mutant clone cells, apt-e2f1-double mutant clone cells, and neighboring control cells. Data were presented as mean ± SD. n = 4 cells for control, 3 cells for e2f1-mutant clone, and 3 cells for apt-e2f1-double mutant clone. ***p < 0.001 versus the control (Student’s t test). See also Table S4.
Apt Up-regulates Rbf1 Also in Mitotic Cycling Cells

So far, we demonstrate the roles for Apt in the induction of CycE and chromatin compaction for silencing in endocycling salivary gland cells. Then, how is the situation in mitotic cycling cells? We have shown that Apt activates the cycE expression for S phase entry in imaginal disc cells undergoing mitotic cycles (Liu et al., 2014; Wang et al., 2017). This led us to examine whether Apt up-regulates Rbf1 also in the wing disc. Strong
Figure 4. Apt Regulates Chromatin Compaction through rbf1 in the Salivary Gland

(A) The ratio of nuclear size to DNA content in apt-mutant clone cells, e2f1-mutant clone cells, or control cells. Data were presented as mean ± SD. n = 10 for control, 4 for apt-mutant clone, and 3 for e2f1-mutant clone. ***p < 0.001 versus the control (Student's t test).

(B) Immunostaining of control or apt-mutant clone cells with the antibody against transcribing RNA polymerase II (H14). An apt-mutant clone cell (marked by yellow dotted line) showed prominent signals of transcribing RNA polymerase II compared with control cells. n = 3 with all samples showing the represented phenotype. Scale bar, 20 μm.

(C) ptc-GAL4-driven RNAi knockdown of rbf1 induces de-compaction of chromatin. Compared with control cells, rbf1-knockdown cells exhibited larger nuclei with de-compacted chromatin. n = 10 with all samples showing the represented phenotype. Scale bars, 20 μm. Graph shows the ratio of nuclear size to DNA content. Data were mean ± SD relative to control. n = 100 for control and 100 for rbf1 RNAi. The control samples were normalized to 1. ***p < 0.001 (Student's t test).

(D) RT-qPCR assays for the expression of dp, rbf, e2f2, and Rbf1-target genes (CG4679, gigas, diap3, lpp) in dpp > GFP control and dpp > GAL4-driven apt-knockdown salivary glands. Data were average ± SD relative to the dpp > GFP mRNA level. dpp > GFP samples were normalized to 1. **p < 0.01, ***p < 0.001 (Student's t test).

(E) The expression of Rbf1 in apt-mutant clone cells. Arrows indicate the decreased expression of Rbf1 in the clone. n = 4 with all samples showing the represented phenotype. Scale bar, 20μm.

(F) Up-regulation of Rbf1 in apt-overexpression cells. Yellow arrowheads indicate y+ flipped out cells expressing actin-GAL4 that drives overexpression of apt. n = 4 with all samples showing the represented phenotype. Scale bar, 20 μm. fb, fat body.

(G and H) Chromatin de-compaction upon ptc-Gal4-driven RNAi of apt was suppressed by simultaneous overexpression of Rbf1. Upper panels are low-magnification images. Lower panels show higher-magnification images of the nuclei. n = 10 with all samples showing the represented phenotype. Scale bars, 20 μm. Graph shows the ratio of nuclear size to DNA content. Data were mean ± SD relative to control. n = 100 for control, 100 for apt RNAi, and 100 for control apt RNAi.
knockdown and overexpression of apt induced cell migration and apoptosis, respectively, in the wing disc, which hampered inspection of the nuclear defects. Therefore, we employed mild knockdown or overexpression of apt that was triggered by heat shock-induced flipping out of y+ from actin > y+>GAL4. RNAi knockdown of apt reduced the Rbf1 expression (Figure 5A) and slightly increased the nuclear size/DNA ratio in wing disc cells (Figure 5B). Conversely, overexpression of Apt enhanced the Rbf1 expression (Figure 5C) and reduced the nuclear size/DNA ratio (Figure 5D). These results indicate that Apt up-regulates Rbf1 to direct proper chromatin compaction in wing disc cells also. Collectively, these data suggest that Apt-mediated CycE induction and chromatin compaction are general mechanisms common to both mitotic cycling and endocycling cells.

**Mutual Activation of FSBP and E2f1, and FSBP-Mediated Chromatin Compaction in Mammalian Cells**

As Apt, E2f1, and Rbf1 are evolutionarily conserved transcription factors, the aforementioned mechanisms could also be conserved in mammalian cells. To test the possibility, we focused on the mammalian homologs of these factors, FSBP, E2f1, and Rb. In mouse NIH3T3 cells, RNAi knockdown of FSBP significantly decreased the expression of E2f1 and CycE homologs (CCNE1 and CCNE2) (Figure 6A). Upon E2f1 knockdown, the expression of FSBP, CCNE1, and CCNE2 were attenuated (Figure 6A). These data indicate mutual activation between FSBP and E2f1 and requirement of FSBP and E2f1 for the expression of CCNEs. In addition, knockdown of FSBP reduced the expression of Rb1, whereas overexpression of FSBP increased the Rb1 expression (Figure 6B). Consistently, the target genes of Rb (CDC6 and DHFR) also showed up- and down-regulation in FSBP-knockdown and FSBP-overexpressing cells, respectively (Figure 6B). These data indicate FSBP-mediated up-regulation of Rb. Furthermore, each FSBP-knockdown cell exhibited a large nucleus with less compact chromatin and a lower Rb protein level than the control cell did (Figures 6C–6E). Taken together, these results demonstrate Apt- and FSBP-mediated conserved mechanisms for CycE induction and chromatin compaction.

**DISCUSSION**

This study revealed Apt-mediated up-regulation of two key players in the cell cycle, CycE and Rbf1. What is the significance of this finding? The positive feedback between apt and e2f1 ensures rapid and robust induction of CycE at S phase entry. Apt also up-regulates Rbf1, but Rbf1 is inactivated through phosphorylation with Cdk2. After initiation of S phase, Rbf1 becomes active through de-phosphorylation and represses cycE (Du et al., 1996; Edgar and Orr-Weaver, 2001; van den Heuvel and Dyson, 2008). Together with Crl4−Cdt2-mediated degradation of E2f1 (Zielke et al., 2011), this leads to a rapid decline of CycE. Therefore, Apt governs both induction and subsequent repression of cycE with the aid of the periodic phosphorylation and de-phosphorylation of Rbf1.

E2f1, a heterodimer of E2f1 and Dp, has been studied for many years, and it is a key regulator of CycE expression for S phase entry (Duronio et al., 1995). However, residual S phase takes place in a null mutant of Drosophila e2f1 or dp (Duronio et al., 1998; Royzman et al., 1997). Here, we solved the discrepancy: another factor Apt also participates in the activation of cycE. Until this study, contribution of “another factor” if any was thought to be rather trivial compared with that of E2f1, because e2f1 or dp mutation severely reduced the CycE expression. Our study revealed that the notion is not correct. As Apt and E2f1 up-regulate each other and both Apt and E2f1 are required for the cycE expression, disruption of e2f1 or dp function leads to depletion of both E2f and Apt, which in turn causes a severe defect in cycE expression. This masked the contribution of “another factor” Apt. Actually, transgenic reporter assays indicated that both the Apt-binding sites and the E2f1-binding sites in the regulatory region of cycE are necessary for the normal level of cycE transcription.

apt-Mutant cells induced in the salivary gland exhibited abnormal nuclei. The size of nucleus/DNA content was ~6.5 times higher than that of control cells, which resulted in de-compacted chromatin. Our study suggests that the unusual phenotype is due to release from Rbf1-mediated chromatin compaction and
Figure 5. Apt Regulates the Expression of Rbf1 and Chromatin Compaction in the Wing Disc
(A) Immunostaining of a wing disc harboring apt-knockdown clones with anti-Rbf1 antibody (red). GFP (green) represents the region of y' flipped out apt-knockdown cells. n = 20 with all samples showing the represented phenotype. Lower panels are close-up images around the apt-knockdown clone. Scale bars, 20 μm.

(B) The wing disc nuclei of control cells (without GFP) and apt-knockdown cells (with GFP) were stained with DAPI. n = 3 with all samples showing the represented phenotype. Red dotted line indicates the boundary between control cells and apt-knockdown cells. Scale bar, 20 μm. Graph shows the ratio of nuclear size to DNA content in the control or apt-RNAi cells. Data were mean ± SD relative to control. n = 55 for control and 32 for apt-RNAi. The control samples were normalized to 1. **p < 0.01 (Student’s t test).
de-repression of Rbf1-target genes that occupy many loci throughout the genome (Korenjak et al., 2012). The nuclear size/DNA content of e2f1-mutant cells was also higher than that of control cells. However, the difference was less prominent than that between apt-mutant cells and control cells. We surmise the following explanation for it. Within a cell, there might exist a balance between the amounts of the E2f1/Dp complex and those of the Rbf1/E2f2/Dp complex. In e2f1-mutant cells, the level of the latter complex would increase in the absence of the former complex. This would compensate the decrease in Rbf1 and E2f2 due to reduced Apt, and would direct toward chromatin compaction.

This study underscores the importance of FSBP, a hitherto not-well-characterized transcription factor. Here we found FSBP- and Apt-mediated up-regulation of Rb and Rbf1, respectively. This raises an intriguing possibility that FSBP (Apt) suppresses tumor metastasis through up-regulation of Rb (Rbf1). Future studies should address the issue experimentally.

Figure 5. Continued
(C) Immunostaining of a wing disc harboring apt-overexpression cells with anti-Rbf1 antibody (red). GFP (green) implies the y′-flipped out apt-overexpressing region. n = 25 with all samples showing the represented phenotype. Scale bar, 20 µm.
(D) The wing disc nuclei of control cells (without GFP) and apt-overexpression cells (with GFP) were stained with DAPI. n = 3 with all samples showing the represented phenotype. Red dotted line indicates the boundary between control cells and apt-overexpression cells. Scale bar, 20 µm. Graph shows the ratio of nuclear size to DNA content. Data were mean ± SD relative to control. n = 40 for control and 52 for apt-overexpression. The control samples were normalized to 1. ***p < 0.001 (Student’s t test).

Figure 6. Mutual Activation between FSBP and E2f1, and FSBP-Mediated Chromatin Compaction in Mouse NIH3T3 Cells
(A) RT-qPCR assays for relative mRNA levels of FSBP, E2f1, CCNE1, and CCNE2 in FSBP-knockdown cells or E2f1-knockdown cells. Data were mean ± SD relative to Mock. *p < 0.05, **p < 0.01 (Student’s t test).
(B) Relative mRNA levels of FSBP, Rb1, CDC6, and DHFR from FSBP-knockdown cells or ectopic FSBP-expressing cells. Data were mean ± SD relative to Mock. *p < 0.05, **p < 0.01, ***p < 0.001 (Student’s t test).
(C) Control and FSBP-knockdown cells were stained with DAPI (blue) and anti-FSBP antibody (green). n = 81 cells for control and 25 cells for FSBP knockdown with all samples showing the represented phenotype. Scale bar, 20 µm.
(D) Control and FSBP-knockdown cells were stained with DAPI (blue) and anti-Rb1 antibody (red). n = 85 cells for control and 26 cells for FSBP knockdown with all samples showing the represented phenotype. Scale bar, 20 µm.
(E) Quantification of the nuclear size. Data were mean ± SD relative to Mock. n = 16 for control and 6 for FSBP RNAi. ***p < 0.001 (Student’s t test).
Limitation of the Study
We demonstrate here Apt-dependent up-regulation of rbf1. There exists a single Apt-binding motif at 156 nucleotides upstream of the transcription start site of rbf1. ChIP assays showed occupancy of Apt on the motif. Therefore, it is most likely that Apt directly activates rbf1 transcription through the binding site. However, further functional analyses including disruption of the Apt-binding site are necessary to verify the possibility.

Resource Availability
Lead Contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Susumu Hirose (shirose@nig.ac.jp).

Material Availability
This study did not generate new materials.

Data and Code Availability
This study did not generate new datasets.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101369.

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AUTHOR CONTRIBUTIONS
X.-F.W., Q.-X.L., S.H., and E.S. designed the project. S.H. and Q.-X.L. supervised the study. X.-F.W. performed the majority of the experiments and analyzed the data with the following exceptions. J.-X.L. and Z.-Z.Z. did the in vitro cell culture experiments and data analysis. Z.-Y.M. helped with the injection to prepare the transgenic flies. Y.S. and H.-R.Z. helped with a part of the genetic and staining experiments. X.-F.W. wrote the original draft. S.H. and Q.-X.L. revised the manuscript.

DECLARATION OF INTERESTS
The authors declare no conflict of interest.

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Supplemental Information

Evolutionarily Conserved Roles for Apontic in Induction and Subsequent Decline of Cyclin E Expression

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Figure S1. Supporting data for mutual activation of apt and e2f1, Related to Figure 1. (A) Expression of dpp-GAL4 in the salivary gland. A dpp>GFP salivary gland stained with DAPI (blue) and anti-GFP antibody (green). (B) Expression of Apt and E2f1 depend on each other. Upper panels: A salivary gland carrying aptRNAi cells was stained with anti-E2f1 antibody. Arrowheads indicate significantly reduced expression of E2f1 in the apt-knockdown cells. Lower panels: A salivary gland carrying e2f1-mutant cells was stained with DAPI and anti-Apt antibody. Arrowheads indicate severely reduced expression of Apt in the e2f1-knockout cells.
Figure S2. Sequence of the *apt* promoter region for reporter assays, Related to Figure 1.
AptPlacZ carries a control sequence with wild-type E2f1-binding sites. AptMPlacZ harbors mutated E2f1-binding sites.
Figure S3. Reporter assays for cycE transcription, Related to Figure 3.

(A) Sequence in the cycE first intron for reporter assays. The original sequence contains a BamH1 enzyme site between the two E2f1-binding motifs. In order to use EcoR I and Bam HI for insertion into the vector, we disrupted the internal BamH1 site by base substitutions. cycEPlacZ is used for control. E2f1BSMPlacZ contains mutated E2f1-binding motifs. E2f1BSM+AptBSMPlacZ contains mutated E2f1- and Apt-binding motifs.

(B) ptc-GAL4-driven RNAi knockdown of apt reduced the LacZ level in E2f1BSMPlacZ. Data were mean ± SD relative to the level of ptc-GAL4 control. n=20 for ptc>aptRNAi and 20 for ptc-GAL4.

***P<0.001.
Figure S4. *cycE* regulatory elements, Related to Figure 3.  
(A) Schematic presentation of the regulatory elements of *cycE*. U element: upstream region of *cycE* TSS. I element: intron (1st) region of *cycE*. TSS: transcription start site.  
(B) The eye disc from *cycEPlacZ* was stained with DAPI (blue), anti-LacZ antibody (green) and anti-CycE antibody (red).
Figure S5. Higher magnification image of apt mutant nuclei, Related to Figure 4. apt mutant nuclei of the salivary gland (without GFP) exhibited intra-chromosomal chromatin de-compaction and large inter-chromosomal spaces compared with control nuclei (with GFP). Dotted line indicates an apt mutant clone.
Table S1. Primers for RT-qPCR, Related to Figures 1, 4 and 6.

apt-F: CGTCTCAGTGTGTCGCCCTAA (Liu et al., PNAS, 2014)
apt-R: CGTGGCGGATATGTTGTCTCA (Liu et al., PNAS, 2014)
e2f1-F: CGCCTCAGTGTGTCGCCCTAA (Zielke et al., Nature, 2011)
e2f1-R: CGCCTCAGTGTGTCGCCCTAA (Zielke et al., Nature, 2011)
cyec-F: GCCATTCTTCCGAGTGTAGCT
ocyec-R: GGCAATAAGCACTTCTGTCA
Dp-F: CCAAGGACAAGAGAAGAATT
Dp-R: ACGATGAGAAGGGAAGCATG
Rbf-F: CGGCAACAAAGGAACATAT
Rbf-R: CATCGTAGGCACACTGAA
E2f2-F: GCTCAACGTGGATCTCTTCA
E2f2-R: CGCCTCAGTGTGTCGCCCTAA
CG4679-F: AGTTCCAGATTTCTGAGATG
CG4679-R: GCAAGTCTGCTACTTCTCTT
Gigas-F: ATAAATGGGTGTCGCCGTCT
Gigas-R: CCTCTGCTCTGTTGCTTG
Diap3-F: CAACGGAGAAAGTACACTT
Diap3-R: CCAGCGAAGCCAGTGA
lpp-F: AGGTTGGATGTGCTACTTT
lpp-R: CTCAATCAGTTACTTCAATG
lacZ-F: CGAAGTGAAGCAAGGAAATA
lacZ-R: GTAGTTCAGGCAGTTCAATC
tubulin 56D-F: GTTGACACTCGGTGTTAGCG (Nishioka et al., 2018, Development)
tubulin 56D-R: CCAAGTGATGCGCTCTGCAATA (Nishioka et al., 2018, Development)
FSBP-F: TTTCGGAGCCCACCAAGCAA
FSBP-R: GCCAGTCTCAGAGCAATTTTGC
CCNE1-F: GTGGCTCCGACCTTTGCT
CCNE1-R: CACAGTCTGCTCAATCTGGGC
CCNE2-F: TCTGTGCATTCTAGCCATT
CCNE2-R: ATCCAGTCTCACACTCCGAG
e2f1-mouse-F: AGAAAACGCGCATCTATGAC
e2f1-mouse-R: CTCAAGCCCGCTTACATC
rb1-F: TCTCACTCCTGACACTT
rb1-R: TCTCACTCCTGACACTT
mcdc6-F: AATGTGAGATCGGATCGTCA
mcdc6-R: AAATTCACCCCTGTTCCCATC
mDHFR-F: CATGGTTTGAGTGCAGGG
mDHFR-R: GTCAAAAAAGATCGGATGG
GAPDH-F: TGTTCCCCTGCGTGGAGTGA
GAPDH-R: TGCTGTGGAAGTCGCAAGGAG
### Table S2. Primers for ChiP, Related to Figures 3 and 4.

| Primer   | Sequence                        | Notes                        |
|----------|---------------------------------|------------------------------|
| E2f1BS-F | GCCACCAGCGGCTTCATCG             |                              |
| E2f1BS-R | CGGCAGGCAAGGATGATC              |                              |
| AptBS1-2-F | CTCGTACAGCTGTTTTTCGAC | AptBS1-2-R: GGCAAGCTTGGCATTTTTATT |
| AptBS3-F | GCCACCAGCGGCTTCATCG (same as E2f1BS) |                              |
| AptBS3-R | CGGCAGGCAAGGATGATC (same as E2f1BS) |                              |
| AptBS4-F | TGATACCCAGTTCACCA               |                              |
| AptBS4-R | CAAAGACACACAGCGGCAA             |                              |

| Primer   | Sequence                        | Notes                        |
|----------|---------------------------------|------------------------------|
| E2f1BS-Ctr1-F | CTTGCGATCATTTGGTTACT         |                              |
| E2f1BS-Ctr1-R | GGACTGGAATGATGGA            |                              |
| E2f1BS-Ctr2-F | AATGAGTGAGCGAGATAGAC       |                              |
| E2f1BS-Ctr2-R | AAGCAGGAGGAAGAGGAA         |                              |
| AptBS-Ctr1-F | AATGAGTGAGCGAGATAGAC       | (same as E2f1BS-Ctr2-F)    |
| AptBS-Ctr1-R | AAGCAGGAGGAAGAGGAA         | (same as E2f1BS-Ctr2-R)    |
| AptBS-Ctr2-F | AACTCCGACTCTCTCTGTGC       |                              |
| AptBS-Ctr2-R | AAGTGGCAGCTGTGAGGAT        |                              |
Table S3. Primers for reporter assays, Related to Figures 1 and 3.

| Promoter          | Forward Primer | Reverse Primer | Restriction Enzyme |
|-------------------|----------------|----------------|--------------------|
| apt promoter      | Reporter-AptF: GGAATTCCTATAATCGCCCATTTAGTC (EcoRI) | Reporter-AptR: CGGGATCCGTTACTGATAAGCTAT (BamHI) | E2F11-AptR: GAAGATCTAGGGCGCTCAACGGCCA E2F11-AptF: GAAGATCTGAACGGAGCGGTATTCG |
|                   | E2F12-AptR: GACTAGTGAGCATGTCGTGCT | E2F12-AptF: GACTAGTGAGCGGGGTCCATACGGA |
| cycE promoter     | Reporter-cycE-F: GGAATTCGTCGAAGTGTTGAA (EcoRI) | Reporter-cycE-R: CGGGATCCATCTTTTGTGACATAGCGCT (BamHI) | BamH1-R1: CGGCTAGCGAAAGCTTTTGCAACACTTTTCGCTCAT |
|                   | BamH1-F1: CGGCTAGCGCTTTTCGCGGCTTTCTCAT | Amt1-R: GACTAGTGGTTGACGTCGATCACG | Amt1-F: GACTAGTGGTTGACGTCGATCACG |
|                   | Amt2-R: CCCTCGAGAGTCCAGTCGCTCAGTCTCAG | Amt2-F: CCCTCGAGAGTCCAGTCGCTCAGTCTCAG |
|                   | E2f11-R: ATTTGCGGCCGCACTGCAACACAGGGG | E2f11-F: ATTTGCGGCCGCACTGCAACACAGGGG |
|                   | E2f12-R: ATTTGCGGCCGCACTGCAACACAGGGG | E2f12-F: ATTTGCGGCCGCACTGCAACACAGGGG |
|                   | Apt3-R: CCCCCGGGTGGAACTGGGAACCTGACGGAACGCGA | Apt3-F: CCCCCGGGTGGAACTGGGAACCTGACGGAACGCGA |
|                   | Apt4-R: GCTCTAGATTTTACCTTCGTTTATTTTAGTGGATA | Apt4-F: GCTCTAGATTTTACCTTCGTTTATTTTAGTGGATA |
Table S4. Example for measurement, Related to Figure 2B

| Round 1 | Area | Mean  | Relative Intensity |
|---------|------|-------|--------------------|
| mutant 1 | 0.034 | 1440.647 | 0.180248168 |
| Ctrl1 for mutant 1 | 0.022 | 6003.591 | 0.751146032 |
| Ctrl2 for mutant 1 | 0.018 | 8622.778 | 1.078848556 |
| Ctrl3 for mutant 1 | 0.018 | 8647 | 1.081879119 |
| Ctrl4 for mutant 1 | 0.02 | 7215.8 | 0.902812923 |
| Ctrl5 for mutant 1 | 0.017 | 9473.706 | 1.185313369 |

average of Round 1 control 7992.575

| Round 2 | Area | Mean  | Relative Intensity |
|---------|------|-------|--------------------|
| mutant 2 | 0.051 | 1126.627 | 0.108180336 |
| mutant 3 | 0.035 | 1464.314 | 0.140605525 |
| mutant 4 | 0.028 | 1865.071 | 0.179086786 |
| Ctrl1 for mutant 234 | 0.021 | 8297.333 | 0.796721786 |
| Ctrl2 for mutant 234 | 0.017 | 10448.06 | 1.003237574 |
| Ctrl3 for mutant 234 | 0.015 | 12161.47 | 1.167761461 |
| Ctrl4 for mutant 234 | 0.015 | 12595.4 | 1.209428329 |
| Ctrl5 for mutant 234 | 0.02 | 8569.45 | 0.82285085 |

average of Round 2 control 10414.34

Control

| Control | 0.751146 |
|---------|----------|
|         | 1.078849 |
|         | 1.081879 |
|         | 0.902813 |
|         | 1.185313 |
|         | 0.796722 |
|         | 1.003238 |
|         | 1.167761 |
|         | 1.209428 |
|         | 0.822851 |

mutant (relative to control)

| mutant | 0.180248 |
|--------|----------|
|        | 0.10818  |
|        | 0.140606 |
|        | 0.179087 |

average of Round 1 control 7992.575

average of Round 2 control 10414.34
Transparent Methods

Fly strains

_Drosophila_ lines were raised in a cornmeal-based regular fly medium (Kayashima et al., 2005). _hsFlp; FRT42D, Ubi-GFP and FRT42D, apt<sup>P₃4</sup> have been described (Wang et al., 2017). _hsFlp; FRT 82B, Ubi-mRFP and FRT 82B lines were gifts from Dr. Tamori. _e2fl_ mutants, _e2f1<sup>07172</sup> (BDSC: 11717) and _e2f1<sup>rm729</sup> (BDSC: 35849), _dpp-GAL4_ (BDSC: 7006), _UAS-GFP_ (BDSC: 35544), _ptc-GAL4_ (BDSC: 81616), _UAS-rbf1_ (BDSC: 50746) and _UAS-rbf1RNAi_ (BDSC: 36744) were obtained from Bloomington Drosophila Stock Center. _UAS-aptRNAi_ (VDRC: v4289) and _UAS-e2f1RNAi_ (NIG-fly: HMS01541) were from Vienna Drosophila Resource Center and Fly Stocks of National Institute of Genetics, respectively. The fly of _FRT 82B, e2f1<sup>07172/TM6B_ was established through recombination. To induce mutant clones of _apt_ or _e2f1_, 2-6 hours AEL embryos were heat shocked at 37 °C for 40 minutes and then reared at 25 °C. _UAS-apt_ was a gift from Dr. Montell. _hsFLP; act>y+>GFP (Ay-GLA4) was a gift from Dr. Tamori and the flip-out cells were induced by a 12-minutes heat shock to second instar larvae.

Immunohistochemistry and Confocal Imaging

Salivary glands or discs were dissected in phosphate buffered saline (PBS), fixed in 4% formaldehyde in PBS for 20 minutes. After washing with PBS with 0.1% Triton X-100, the samples were stained with antibodies in the same solution. We incubated samples with primary antibodies at 4°C for overnight with shaking, and then washed samples for 15 minutes three times. Primary antibodies were used at following dilutions: Rabbit
anti-Apt (1:1000); goat anti-E2f1 (1:100, Aviva Systems Biology, OAEB03032); goat anti-CycE (1:200, Santa Cruz, sc-15903); mouse anti-LacZ (1:500, DSHB, 40-1a); mouse anti-Rbf1 (1:50, a gift from Dr. Dyson); mouse anti-H14 (1:100, Covance, 920304); rabbit anti-FSBP (1:500, ATLAS ANTIBODIES, HPA025059); rabbit anti-Rb1 (1:200, ABclonal, A11409). The secondary antibodies conjugated with Alexa 488 (Molecular Probes, R37114), Cy3 (Jackson ImmunoResearch, 711-165-152, 715-165-150, 705-167-003) and Cy5 (Jackson ImmunoResearch, 711-175-152, 715-175-150) were diluted 1:500 and incubated at room temperature for 2 hours. After washing, samples were mounted and imaged with an Olympus FV1200 Confocal Microscope.

**Image analysis**

For nuclear size and DNA content analysis, images from Confocal were quantified using Analyze function of the ImageJ software according to the reference (Zielke, et al., 2011). For the measurement of clone cells and their neighboring controls, we did z-stack using confocal sections. Nuclei that visually overlapped with their neighboring cells were not analyzed. Regions were selected using freeform selection tool. The area function was used to measure nuclear area. DNA content was quantified by DAPI mean grey value. An example of the measurement was shown in Table S4. Statistical analyses on LacZ expression, DNA content, nuclear size and nuclear size / DNA content data were carried out with Student’s $t$-test.
Motif search

The regulatory sequences of *apt* and *cycE* were obtained from UCSC (http://genome.ucsc.edu). E2f1- and Apt-motifs were derived from published references (Kel et al., 2001; Korenjak et al., 2012; Liu et al., 2003; Yamaguchi et al., 1995). E2f1-binding motif used in this study was TTTGGCGC or CTTCGCGG. Apt-binding motif was (T)CCAATT(G).

Chromatin immunoprecipitation (ChIP)

ChIP experiments were performed as previously described (Wang et al., 2017). Briefly, 200 early third instar salivary glands were dissected, fixed, homogenized and sonicated. After analyzed the fragment sizes, sonicated samples were immunoprecipitated with antibodies. 2μl of purified DNA from ChIP samples and input were amplified using specific primers (Table S2). Statistical analyses on ChIP data were carried out with Student’s *t*-test.

Plasmid constructions and transgenic flies

*AptPlacZ* was made by inserting a 1.5 kb *apt*-promoter region that contains E2f1-binding motifs into a CaSpeR-AUG-β-gal vector (Thummel et al., 1988). AptMPPlacz with mutated E2f1-binding motifs was made from *AptPlacZ*. *cycEPlacZ* construct was made by inserting a 3 kb region in the first intron of *cycE* carrying Apt- and E2f1-binding motifs into a hsCaspeR-AUG-β-gal vector (Thummel et al., 1988). *E2fBSMPlacZ* with mutated E2f1-binding motifs and *E2fBSM+AptBSMPlacZ* with
mutated E2f1- and Apt-binding motifs were made from *cycEPlacZ*. Primers used for mutagenesis of the binding motifs are shown in Table S3. To generate transgenic fly lines, constructs were injected into the germ line. After chromosomal mapping, flies with normal *mini-w*⁺ expression were used for reporter assays.

**RT-qPCR**

Total RNAs were prepared from 40 dissected early third instar salivary glands using an RNAprep kit (Zymo Research, Cat. number R2050). cDNAs were synthesized from RNA samples of three biological replicates. qPCR was performed as described previously (Wang et al., 2017). Primer sequences used for qPCR are shown in Table S1. The amount of mRNA was normalized to that of β-tubulin mRNA and then presented as fold change against the control mRNA level. Statistical analyses were performed by Student’s *t*-test.

**Cell culture and transfection**

NIH3T3 cells were grown in Dulbecco’s modified Eagle medium (Biological Industries, REF number 06-1055-57-1A) supplemented with 10% Fetal Bovine Serum (FBS) and 1% penicillin/streptomycin. To silence FSBP or E2f1, siRNAs were transfected at a final concentration of 100nM using lipofectamine 2000 (Invitrogen, Product number 11668019) according to the manufacturer’s instruction. The siRNA sequences used were: Negative control: 5’-UUCUCCGAACGUGACGUCUdTdT; FSBP-siRNA: 5’-GCAAGUCAUGGAAUUGAUUTdTdT; E2F1-siRNA:
5’-GGAUCUGGAGACUGACCAUTTdTdT. To generate Fg-mFSBP for mammalian cell expression, we amplified this gene via cDNA and cloned it into CMV-Fg constructs. The primer pairs used were as follows: mFSBP-Forward: 5’-ATGGTAGGAAAGGCTAGATC-3’ and mFSBP-Reverse: 5’-TCAGAGACTACTGTATTGAG-3’. Cells were harvested for RT-qPCR analyses as previously described (Wang et al., 2017). Primers used were shown in Table S1.

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