Endocycles support tissue growth and regeneration of the adult Drosophila accessory gland

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Abstract

The *Drosophila melanogaster* accessory gland is a functional analog of the mammalian prostate made up of two secretory epithelial cell types, termed main and secondary cells. This tissue is responsible for making and secreting seminal fluid proteins and other molecules that contribute to successful reproduction. Here we show that similar to the mammalian prostate, this tissue grows with age. We find that the adult accessory gland grows in part via endocycles to increase DNA content and cell size, independent of mating status. The differentiated, bi-nucleated main cells remain poised to endocycle in the adult gland and upregulation of signals that promote endocycling and tissue growth are sufficient to trigger dramatic endocycling leading to increases in cell size and ploidy. The main cells of this tissue remain poised to enter the cell cycle and endocycling of main cells increases during recovery from severe tissue damage. Our data establish that the adult accessory gland is not quiescent, but instead uses endocycles to maintain the accessory gland’s critical function throughout the fruit fly’s lifespan.
Introduction

The *Drosophila* accessory gland is functionally analogous to the mammalian prostate. This tissue is an essential component of the male reproductive system and is responsible for making and secreting seminal fluid proteins, sex peptides, and antimicrobial proteins that are transferred to the female upon mating (Ravi Ram, Ji, and Wolfner 2005; Qazi and Wolfner 2003; Lung, Kuo, and Wolfner 2001; Adams and Wolfner 2007; Heifetz et al. 2005). The accessory gland is comprised of a single layer of secretory epithelial cells that form a large lumen apically and are surrounded basally by extracellular matrix and a muscle layer (Susic-Jung et al. 2012; Bairati 1968). Each lobe of the accessory gland consists of approximately 1,000 epithelial cells, which are made up of two cell types: main cells and secondary cells (Bairati 1968). Main cells are the smaller of the two types and are hexagonal in shape. These cells make up a majority of the gland and are located mostly in the proximal and medial portions of the lobes. Secondary cells are larger, more luminal cells that are located at the distal tip of the lobes. There are approximately 40-60 secondary cells in each lobe, with the rest of the cells (~940-960 cells) being main cells. Main and secondary cells have distinct but partially overlapping secretory protein profiles, and both cell types play an important role in fecundity (Bertram et al. 1992; Sitnik et al. 2016).

In late larval stages, FGF signaling drives recruitment of mesodermal cells to the genital disc. These cells undergo a mesenchymal to epithelial transition and give rise to the precursors for the accessory glands and seminal vesicles (Ahmad and Baker 2002). During early metamorphosis, the accessory gland progenitors increase in number by standard mitotic cell cycles. Approximately 50-55 hours after pupa formation, the cells of the developing accessory gland arrest proliferation and synchronously enter a truncated variant cell cycle, in which nuclear division occurs but cytokinesis does not, resulting in the bi-nucleation of the epithelial cells of the accessory gland. Approximately 10 hours later, the cells enter an additional synchronized endocycle, increasing their DNA content without mitosis. This results in ~1,000 binucleate cells containing two 4N nuclei. After this endocycle, the cells of the accessory gland exit the cell cycle (Taniguchi et al. 2014) and are thought to remain quiescent in the adult (Leiblich et al. 2012), with the exception of
secondary cells, which retain the capacity to cycle in response hormonal or mating signals (Leiblich et al. 2019).

We examined endoreplication in the adult accessory gland from eclosion and find many examples of endoreplication in main cells, independent of mating status. Induction of cell cycle and growth regulators are sufficient to trigger a dramatic increase of endocycling in the differentiated adult tissue and main cells, even in the absence of mating. Importantly, we find that main cells remain poised to re-enter the cell cycle to maintain the capacity for gland regeneration upon tissue damage, regardless of mating status. Our work establishes that adult accessory gland main cells are binucleate, polyploid cells that remain poised to endocycle to maintain gland function throughout the fruit fly's lifespan.
The adult accessory gland grows via endoreplication, independent of mating status, under normal physiological conditions throughout lifespan.

Like the mammalian prostate, the Drosophila accessory gland (AG) grows with age. AGs from virgin males undergo a period of rapid growth from the day of eclosion to day 10 post eclosion (Fig 1A), and continues to grow, albeit at a lesser rate throughout adulthood (Fig 1B). While some AG growth in early adulthood can be attributed to the production and secretion of sex peptides which expands the lumen of the gland, we hypothesized that changes in cell number or cell size may also contribute to gland growth. To address whether a mitotic population exists in the AG, we stained the adult AG for the mitotic marker, phospho-histone H3 (PH3), at various timepoints throughout the male lifespan. In sum, over 100 AGs, of various ages with and without mating, were negative for PH3 staining (data not shown). We therefore postulated that under normal physiological conditions, gland growth is due to changes in cell size rather than cell number.

We next measured cell size for the major population of cells in the AG, the main cells. Antibody staining for the septate junction protein Discs large (Dlg) was used for all cell size measurements. Because of the apical localization of Dlg, cell size measurements reported here are not the full volume of a cell, but rather a measurement of the apical area. In virgin males, main cell size increases with age throughout lifespan (Fig 1C), and the variation in main cell size throughout the gland also increases with age. Previous studies have established that cell size often scales with nuclear size (Orr-Weaver 2015). We therefore next measured nuclear size in main cells across lifespan and found that nuclear area and variance also tracks with the increases in main cell size with age (FIG 1C). Together, our data shows a correlation between increased AG size, main cell size, and main cell nuclear area throughout the lifespan of the fly (Fig 1).

Many tissues undergo a variant cell cycle called an endocycle in order to increase tissue size under normal physiological conditions (Orr-Weaver 2015; Edgar, Zielke, and Gutierrez 2014). Endocycling occurs when cells cycle through G/S phases without entering an M phase, thereby increasing DNA content resulting in polyploidy. Increased ploidy can contribute to increased biosynthesis, as is the case with Drosophila nurse cells (Orr-Weaver 2015; Inge Øvrebø and Edgar 2018). In this context, endocycling may be essential for optimal function of the adult AG, which is highly secretory and responsible for making large
amounts of AG specific proteins and other molecules for transfer to the female upon mating (Prince et al. 2019; Wilson et al. 2017). Because cell size scales with DNA content in Drosophila endocycling cells (Von Stetina et al. 2018), we next examined whether DNA replication occurs in the main cells of this tissue. We first examined the AG immediately after eclosion, before the males eat or mate. To label S-phases we incubated tissues with EdU for one hour, since we found labeling by feeding to be ineffective at this stage. Upon a one-hour EdU incubation, we observed extensive nuclear labeling in most main cells, demonstrating that the adult AG undergoes a very early wave of endocycling in main cells (Fig 1D, top row). We hypothesize that the burst of widespread EdU incorporation that we see results from a relatively synchronous round of endoreplication occurring within the first few hours after eclosion that likely supports the extensive growth and maturation of this gland during the first few days of adulthood (Koppik et al. 2018). We believe this round of endoreplication in the AG has been previously missed due to the use of feeding-based labeling assays, since adult male flies do not eat for 4.5-5 hours post eclosion.

To assess endocycling in aging adults, we performed EdU incorporation experiments by feeding adult male flies an EdU/sucrose mixture for 10 day windows at different time points throughout their lifespan. This longer-term EdU labeling allowed us to visualize slow or rare DNA replication events in the adult AG. We found that the main cells of the AG continue to endocycle throughout the lifespan of the fly up to 50 days of age (Fig 1D) under both starved and fed conditions (fed data not shown). We only very rarely observed secondary cells labeled with EdU, and this was independent of mating status (Fig 1E,F). We conclude that adult AG main cells undergo endocycling throughout the Drosophila lifespan under normal, physiological conditions with and without mating.

**Cell cycle degradation machinery oscillates in the adult accessory gland**

Our pulsed and long-term EdU labeling approach provides a static picture of S-phase but does not reveal cell cycle transitions and oscillations in real-time. To address this, we used Fly FUCCI cell cycle reporter to examine the oscillations of the cell cycle degradation machinery in the AG (Zielke et al. 2014). We used Paired-Gal4 (Prd-Gal4), a driver that has been established to express in both main and secondary cells, to drive expression of the cell cycle protein degrons mRFP1-CycB1–266, to assay APC/C[Cdh1] activity, and GFP-E2F11–230, to.
assay CRL4(Cdt2) activity. (Fig 2A). We find that more than 24h after eclosion, the majority of main and secondary cells have low levels of mRFP1-CycB1-266 indicating most cells are normally in a Gap phase with high APC/C<sup>Cdh1</sup> activity, except for a small subset of main and very rare secondary cells where high mRFP1-CycB1-266 indicates low APC/C<sup>Cdh1</sup> activity. The low APC/C<sup>Cdh1</sup> activity in endocycles results from transient high CycE/Cdk2 which also triggers S-phase, suggesting these cells have recently undergone DNA synthesis (Narbonne-Reveau et al. 2008; Zielke et al. 2008). The relatively low numbers of these cells at individual fixed timepoints roughly agrees with the number of S-phases we observe in our longer term EdU labeling. The GFP-E2F1<sub>1-230</sub> reporter indicates CRL4(Cdt2) activity which is normally high during S-phase and low during Gap phase. The high levels of GFP-E2F1<sub>1-230</sub> throughout the gland verifies that most cells are in a Gap phase where CRL4(Cdt2) activity is low, but a number of main cells and rare secondary cells show high CRL4(Cdt2) activity, consistent with transitions through S-phase. This Fly FUCCI signature is similar to other endocycling cells, such as the intestinal enterocytes (Zielke et al. 2014).

We verified the Fly FUCCI results by using the RGB cell cycle sensor as an additional tool to track cell cycle degradation machinery activity (Fig 2B). The RGB cell cycle sensor uses the Gal4-UAS system to drive overexpression of cell cycle protein degrons nlsCycB<sup>1-96</sup>-nlsCycB<sup>1-285</sup>-tdTomato, to assay APC/C activity, and nlsCdt1<sup>1-101</sup>-EBFP2, to assay CRL4(Cdt2) activity, as well as a full length EGFP-PCNA to visualize early S Phase (Handke et al. 2014). While similar in concept to Fly FUCCI, a few notable differences include: a different length fragment of CycB, the use of nlsCdt1<sup>1-101</sup> instead of E2F1<sub>1-230</sub>, and the expression of all 3 fluorescent reporters from a single transcript. Consistent with our findings using Fly FUCCI, we see that generally APC/C activity is high and CRL4(Cdt2) activity is low throughout the gland with only a small subset of main cells exhibiting EGFP-PCNA foci indicative of S-Phase. Importantly, oscillations of the cell cycle degrons occur at the distal tip where main and secondary cells are located, as well as mid-lobe where there are no secondary cells (Fig 2B).

One caveat of the cell cycle reporters is their dependence on the Gal4-UAS system for ectopic expression of fluorescently tagged degrons. We therefore confirmed our observations using an endogenously tagged PCNA (PCNA::GFP) to confirm the presence of PCNA protein and to use GFP expression as a real-time read out of S-phase activity in this
tissue (Blythe and Wieschaus 2016). PCNA::GFP is present in nearly every main cell on the day of eclosion, the timepoint that we report widespread EdU labeling (Fig 2C, top panel). Consistent with our EdU labeling on the day of eclosion and fly Fucci data, we see lower levels of PCNA::GFP in secondary cells than main cells, suggesting that these cells are refractory to S-phase during this early wave of endocycling. In older AGs PCNA::GFP expression in main cells is greatly reduced (Fig 2C, bottom panel), however we find a small subset of main cells that are PCNA::GFP positive and we did not observe any endogenous PCNA::GFP expression in secondary cells. We conclude that upon eclosion, a wave of endocycling occurs in most, if not all, main cells of the AG leading to dramatic growth and an increase in main cell ploidy in the first day. After that, we find low, but persistent endocycling in several main cells throughout the gland, potentially to maintain tissue growth throughout adulthood, accompanied by very rare (only 2 observed) endocycles in secondary cells.

**Main cells of the adult accessory gland are poised to endocycle**

Our EdU labeling data and cell cycle reporter analysis show that the main cells of the adult AG continue to endocycle throughout the fly’s lifespan. We therefore wondered if these cells remain poised to endocycle in response to cell cycle or cell growth signals.

We used the FLP-FRT system to create clones that express cell cycle and growth regulators in the adult AG (Fig 3A). In brief, this method uses a hs-flp to recombine FRT sites in an actin promoter-driven “flipout” (act-FRT-stop-FRT-Gal4) cassette. Upon removal of the stop codon, the actin promoter leads to constitutive expression of Gal4 and permanent induction of the UAS transgenes. Using this method we titrated the heat-shocks to control the number of cells that express Gal4 and induced subsets of cells in the adult AG to overexpress E2F1 and DP, which are essential for endocycling (Zielke et al. 2011). Ectopic E2F1/DP led to larger nuclei, indicative of extra endocycles (Fig 3B).

We next examined whether overexpression of E2F1/DP with other known endocycle growth regulators could increase adult AG endocycling. *Drosophila Myc* (dMyc) has been shown to play an important role in endocycling in other *Drosophila* tissues, such as the larval salivary gland and the ovary (Pierce et al. 2004; Maines et al. 2004), while the BMP type I receptor Thickveins has been shown to promote growth and endocycling in the
AG secondary cells (Leiblich et al. 2012, 2019). We induced overexpression of dMyc +E2F1/DP, or a constitutively active form of Thickveins, Tkv* + E2F1/DP in a subset of cells in the adult AG. These growth regulators led to dramatic increases in nuclear size and DNA content, easily observable by DAPI staining in the adult AG (Fig 3C). To confirm that the ectopic S-phases occur in the adult, we performed a one day EdU/sucrose feeding on animals at 4 days post eclosion with expression of dMyc +E2F1/DP and observed extensive EdU incorporation in adult tissues as compared to the control animals (Fig 3D). The ease with which the main cells of the AG can be induced to endocycle suggests that the main cells of the AG remain poised to enter the cell cycle upon receiving proper cues.

**dMyc is required for adult accessory gland endocycling and function**

We next asked if dMyc is required for normal levels of gland endocycling and overall gland function. We used 3 dMyc RNAi lines: BL25783 (dMyc-i #1), BL25784 (dMyc-i #2), and BL51454 (dMyc-i #3) driven by Prd-Gal4 to knockdown dMyc in the adult AG. We performed sperm competition assays to test for gland function and EdU assays for endocycling.

To assess sperm competition, we crossed red-eyed males with AGs depleted for dMyc with white-eyed females that were pre-mated to white-eyed males. We then used eye color to track the percentages of F1 progeny that arise from the pre-matings versus second mating to animals with reduced dMyc. During the first 3 days of mating, we saw no progeny from lines with reduced dMyc, compared to 15-30% of progeny coming from second matings in a control without RNAi (Fig 4A). As time progressed, sperm from dMyc-i #1 remained unable to compete, while the other two knockdowns show an increase in sperm competition, eventually reaching normal levels after a delay (Fig 4A). This suggests gland function is disrupted by knockdown of dMyc.

Overall gland size is reduced when dMyc is knocked down (Fig 4B), and nuclear area is reduced (Fig 4C). To confirm that the reduced size is caused by reduced endocycling we measured EdU incorporation and found that that all AGs have reduced EdU incorporation within hours of eclosion (Fig 4D) and with a 10 day feeding (Fig 4E). This demonstrates that dMyc plays an important role in adult AG endocycling, which is critical for proper gland function.
The adult accessory gland can regenerate

We show that the adult AG exhibits a low level of endocycling for normal tissue growth and that this tissue is poised to endocycle. Due to the importance of the AG for successful reproduction, we wondered whether this tissue remains poised to cycle in order to respond to damage.

We used the DEMISE system to induce cell death in the AG. The DEMISE system uses a heat-shock-flippase (hs-Flp) with a UAS-driven "flipout cassette" to induce controlled Reaper expression, driving caspase dependent apoptosis (Cohen et al. 2018). We find that the DEMISE system induces apoptosis in the AG and can be used to study the response to damage in the adult AG (Fig 5). We used three experimental procedures with DEMISE to induce damage and allow for varying levels of recovery (Fig 5A). While DEMISE contains a hs-Flp to induce damage, we quickly realized that all versions of the system we tested lead to leaky expression of the flp enzyme in the late pupal AG after 60h APF at room temperature. Therefore, all DEMISE experiments described in this paper use the leaky expression at late pupal stages to induce damage. When we dissect AGs expressing DEMISE on the day of eclosion (red), we find that 100% of AGs show damage in at least one lobe with the majority of samples exhibiting one lobe with more damage than the other (Fig 5F). When we dissect AGs expressing DEMISE three days post eclosion (blue), we observe that AGs have increased in size since the day of eclosion, but the tissue shows signs of continued damage (Fig 5C), with an increased percentage of animals exhibiting high levels of damage in both lobes (Fig 5F). In a third procedure, we collected animals the day of eclosion, when 100% of animals we tested exhibit damage, and immediately shifted them to 18°C for 5 days to reduce the Gal4/UAS-driven reaper expression to allow for some level of recovery. Indeed, using this protocol we observed striking recovery of gland morphology and size. By 5 days of recovery only 20% of animals show obvious AG defects, compared to 100% of animals that show damage on the day of eclosion (Fig 5D orange outline, Fig 5F).

Importantly, we observed no animal lethality with these DEMISE protocols, so we are confident we are not selecting for survivors at day 5. In all experimental procedures the adult AG contained binucleated, pyknotic nuclei, indicating that Reaper induces apoptosis in AG cells after the binucleation event during development. Further, we observe
binucleated pyknotic nuclei in all of our experiments, regardless of age, suggesting that Reaper continues to induce cell death in the differentiated binucleate adult tissue (Fig 5E), although at reduced levels in our recovery protocol.

We next tested if the recovery that we observed at 18°C restores gland function. We assayed fertility of males with DEMISE active at 25°C and post 5 day recovery at 18°C. When DEMISE is active, AG size and fertility are reduced (Fig 5G left, H), but after recovery tissue size and function are largely restored (Fig 5G right, H), indicating tissue regeneration occurred. Importantly, recovery allows for restoration of tissue size even without mating, indicating that the AG can grow independent of mating status.

**The adult accessory gland exhibits compensatory cellular hypertrophy**

Upon injury, tissues that have a mitotic population increase cell division to allow for regeneration after damage; however, we do not observe an obvious mitotic population in the adult AG. One mechanism of tissue repair in post-mitotic tissues is compensatory cellular hypertrophy (CCH), an increase in cell size that stems from increased endoreplication (Tamori and Deng 2013a). Similarly other postmitotic tissues exhibit wound induced polyploidization (WIP), which can be a result of cell fusion, but is also driven in part by endocycling (Losick, Fox, and Spradling 2013; Losick, Jun, and Spradling 2016). Because the adult AG maintains the capacity to endocycle, we examined whether this tissue may exhibit features of CCH and/or WIP during regeneration.

After recovery from damage, AG lobes with higher levels of pyknotic nuclei exhibit a greater variation in cell and nuclear size, consistent with CCH (Fig 6A,B). However, we never observe cells with more than two nuclei under any protocol, suggesting the cell fusion aspect of the WIP response is absent. We next tested whether recovery from damage leads to an increase in endocycling, as is described in previous work on other post-mitotic tissues (Losick, Fox, and Spradling 2013; Tamori and Deng 2013b). We coupled DEMISE induced damage with a longer recovery including EdU feeding to label cells that endocycle during recovery. Animals were raised at room temperature, collected the day of eclosion, and shifted to 18°C for 11 days and fed EdU/sucrose for the entire recovery (Fig 6C red). We observed increased levels of endocycling throughout the tissue (Fig 6D,F), to an extent much higher than that observed under normal physiological conditions. To address
whether there is a critical time window for damage induced endocycling, we performed the same recovery protocol except that animals were fed standard CM for the first 5 days, and only fed EdU/sucrose for the remaining 6 days (Fig 6C blue). Again, we observed increased endocycling throughout the gland even after 5 days of recovery, demonstrating that increased endocycling continues and is not limited to an early acute damage response (Fig 6E,F). Together, our data suggests that the adult AG is not quiescent, but instead, uses endocycles for normal tissue growth and is poised to endocycle in response to tissue damage. We conclude that the adult AG uses CCH to regenerate from large-scale tissue damage.

**Additional developmental control of variant cell cycles in the adult accessory gland**

The FLP-FRT background we used to induce E2F1/DP and dMyc expression (Fig 3A) also contains a UAS-nuclear GFP (nlsGFP) to visualize which cells express Gal4. As described above, cells that are GFP positive and express dMyc+E2F1/DP exhibit enlarged nuclei through endoreplication (Fig 3C, Fig 7A). However, we noted that neighboring cells that appeared to be GFP negative also often exhibited an enlarged phenotype and endoreplication (Fig 7A). This led us to speculate that these cells may contain ring canals that allow neighboring cells to communicate growth signals in the adult AG.

Ring canals are created when cytokinesis is truncated and the cytokinetic furrow does not fully close and is instead stabilized. The resulting actin-rich structure creates an opening between cells through which cytoplasm is shared (McLean and Cooley 2014). Some, but not all cytoplasmic molecules can travel through these structures, and in particular nlsGFP travels through ring canals at a slow rate (McLean and Cooley 2014). We examined the localization of Pavarotti (Pav), a protein known to localize to ring canals in other *Drosophila* tissues. Using Pav tagged with GFP (Pav::GFP), we and others (Eikenes et al. 2013) observe localization consistent with the formation of ring canals in the adult AG (Fig 7C, top panel). We suggest that the non-autonomous growth effects we observe with dMyc+E2F1/DP are due to dMyc and/or E2F1 complexes passing through AG ring canals. Consistent with this idea, when we carefully quantify GFP fluorescence intensity, we find very low levels of GFP present in the immediate neighbor to the GFP positive cells (Fig 7B).
The location and number of ring canals in the adult AG displayed a distinct and reproducible pattern. Only one ring canal is present on each main cell and is located centrally on membranes at bicellular junctions. Two neighboring main cells can have a single ring canal between them, but no other ring canals between themselves and any other cells. This patterning suggests that ring canals are remnants of a truncated, penultimate cell cycle, generating sister cells prior to the binucleation event during AG development (Fig 7C, bottom panel). Importantly, Pav::GFP is not seen on the membrane of secondary cells, suggesting that these cells do not communicate via shared cytoplasm with their neighboring main cells. This supports a model for the developmentally regulated variant cell cycles of the Drosophila AG main cells that involves a progressive truncation of the canonical mitotic cell cycle. At 40-50 hrs APF we suggest a penultimate cell cycle occurs with a partially truncated cytokinesis to form ring canals in sister main cells. Around 50-60 hours APF, main cells undergo a further truncated cycle in which nuclear mitosis proceeds but cytokinesis does not occur at all, leading to bi-nucleation (Taniguchi et al. 2014). Then at 70-80h APF, the final cell cycle during metamorphosis is an endocyte completely lacking mitosis. The next endocyte begins within the first 5 hours of eclosion, prior to reproductive maturity to increase gland size. During adulthood the mature gland maintains the ability to endocyte to ensure gland function and size is maintained throughout the lifespan of the fly (Fig 7D).
**Discussion**

We find that under normal physiological conditions, cells of the adult AG endocycle and that this contributes to organ growth with age. Importantly, we show that this endocycling occurs both with and without mating and is maintained primarily in the main cell population. We used cell cycle reporters to verify hallmarks of the endocycle such as oscillations of the cell cycle degradation machinery. We employed expression of factors known to regulate endocycling, E2F1/DP, dMyc, and Tkv*, to trigger dramatic increases in endocycling. Finally, we show that this tissue is likely poised to endocycle to maintain proper tissue function throughout the lifespan and to allow for tissue regeneration upon damage. This work establishes that main cells of the adult AG are not quiescent as previously suggested (Leiblich et al. 2012, 2019), but instead exhibit low levels of endocycling normally and remain poised to endocycle in response to specific signals in order to maintain AG function.

**Control of variant cell cycles during AG development**

Our data has revealed additional variant cell cycles during AG development. We show evidence that a penultimate cell cycle with a truncated cytokinesis occurs in the developing gland during metamorphosis to form ring canals and we find a very early wave of endocycling that occurs in the adult AG within hours of eclosion, independent of feeding or exposure to females. Our results reveal a more complete picture of cell cycle regulation in the AG, where the cell cycle is progressively truncated during the later stages of metamorphosis resulting in a binucleate and polyploid tissue (Fig. 7D). Polyploidy further increases during adulthood through additional endocycles under normal tissue homeostasis. This demonstrates the AG is an excellent model system for studying multiple types of variant cell cycles in a developmental context.

**Endocycling during gland maturation and tissue homeostasis**

We describe a wave of endocycling that occurs in most, if not all, main cells of the AG, just after eclosion. We show this wave of endocycling and the resulting early gland growth is important for gland maturation and function. While we show that dMyc plays a role in early gland growth, the identity of the signaling pathways that induce this early wave of
endocycling is not known and will be important for understanding AG maturation. Interestingly, a previous report found that secondary cell nuclei are smaller than main cell nuclei on the day of eclosion, prior to mating (Leiblich et al. 2012). This is consistent with our EdU and PCNA::GFP assays which show that this wave of endoreplication appears to be main cell-specific.

Using a time course of EdU labeling techniques and cell cycle reporter lines, we show that main cells of the AG remain poised to endocycle throughout the fly's lifespan. These cells appear to follow a canonical endocycle, and upregulation of factors previously implicated in growth-dependent endocycling mechanisms, E2F1/DP, dMyc, and Tkv*, are sufficient to induce increased levels of endocycling. We show that dMyc plays a critical role in adult AG endocycling and that normal levels of endocycling are critical for proper tissue function.

**Endocycles in main vs. secondary cells**

Under the conditions examined here, we find that secondary cell endocycles are very rare. This is supported by the patterns of oscillations we observe with cell cycle reporters, which suggest secondary cells are more refractory to endocycling than main cells. Our observations are in contrast to a recent paper that shows secondary cells endocycle in a mating dependent manner and reports a lack of main cell endocycling (Leiblich et al. 2019). There are a few possible reasons as to why we see such dramatic differences in levels and locations of AG endocycling. First, we used different genetic backgrounds in our studies. We examined the Canton-S strain as our wild-type control and used w^{118} strains expressing a paired-Gal4 transgene for our cell cycle reporter studies. Second, our mating protocols may differ. We performed mating under our normal crossing conditions (1:1.5 male to female ratio), rather than multiply-mated or mating-to-exhaustion protocols (Leiblich et al. 2012). Finally, there may be subtle differences in our culturing conditions that lead to differences in AG tissue homeostasis.

Our results using cell cycle reporter lines also hint at a possible mechanism for the differences we observe between main and secondary cell endocycles. We find that GFP-E2F1_{1-230} appears stabilized in secondary cells, consistent with a lack of S-phase coupled degradation. By contrast, nlsCdt1^{1-101}-EBFP2 levels are dramatically lower in secondary
cells than main cells (Supp Fig 1A). This is unexpected since in *Drosophila* the same CRL4(Cdt2) S-phase-dependent degradation is thought to be the major pathway for both CDT1 and E2F destruction (Lee et al. 2010; Zielke et al. 2014). We suggest the nlsCdt1<sup>1-101</sup>-EBFP2 may be degraded by another S-phase independent pathway in the secondary cells of the AG. Mammalian Cdt1 is also degraded via a Cdk2-regulated phosphodegron recognized by the SCF<sup>Skp2</sup> pathway, but this pathway has been thought to play a minor role for *Drosophila* Cdt1 (Zielke et al. 2014). In flies the SCF<sup>Skp2</sup> binding motif RRL contains a substitution to ARL, and the major Cdk2 phosphosite (T29/P in mammals) is not fully conserved. However, there is abundant evidence that fly Cdt1 is phosphorylated by CycE/Cdk2 on multiple sites (Thomer et al. 2004), including on a nearby site conserved in mammals (S31/P). Furthermore, an alternate potential SCF<sup>Skp2</sup> binding RRL motif is found more N-terminally in fly Cdt1 (Supp. Fig 1B), and some genetic evidence suggests fly Cdt1 can also be regulated by the SCF<sup>Skp2</sup> complex in specific cell types (Kroeger et al., 2013). One of the two mammalian Cdk phosphosites in the more recently identified CDT1 PEST domain sequence are conserved in flies and present in the CDT1-BFP construct, but was not interrogated in a previous study of phosphomutant Cdt1 (Thomer et al. 2004; Zielke et al. 2014). We therefore suggest the CDT1-BFP construct could reflect more elaborate Cdt1 regulation than just CRL4(Cdt2) S-phase degradation. Altogether our data suggests secondary cells exhibit rare endocycles under normal physiological conditions, possibly due to specific regulation of replication licensing through secondary-cell specific Cdt1 degradation. This may allow secondary cells to remain poised for endocyte entry upon specific signals.

**The adult accessory gland as model for tissue regeneration**

We show that the adult AG retains the capacity to undergo growth and regeneration upon tissue damage. The tissue exhibits hallmarks of compensatory cellular hypertrophy (CCH) in response to damage, including increased cell and nuclear size along with increased EdU labeling. We hypothesize that canonical endocyte regulators may play a key function in this tissue to mediate CCH and WIP for gland recovery. Importantly, the mammalian prostate also shows regenerative capacity upon partial prostatectomy (Isaacs 2008). The
work here establishes the *Drosophila* AG may be an excellent model for prostate regeneration.
Fly stocks:
Canton S
W^1118
Fly Fucci (BL#55122) w[1118]; Kr[If-1]/CyO, P{ry+[t7.2]=en1}wg[en11]; P{w+[mC]=UAS-GFP.E2f1.1-230}26 P{w+[mC]=UAS-mRFP1.NLS.CycB.1-266}17/TM6B, Tb
RGB Cell Cycle Sensor - P{w+(+mC)=UAS-nlsCdt1N101EBFP2-T2A-nlsCycBN96-nlsCycBN285tdTomato-T2A-EGFPCyO} II.1 (UAS-RGB cell cycle tracker). (Handke et al., 2014)
EGFP::PCNA (provided by S. Blythe)
DEMISE lines: all data shown is with line 10-3; pUAST-FRT-Stop-FRT-rpr/CyO – (provided by D. Fox Lab)
y,w,hs-flp; ; Prd-gal4/TM6B (y,w,hs-flp^12 and BL1947)
w; UAS-P35/Cyo-GFP; act>CD2>gal4,UAS-GFP(nls)/TM3-Ser-GFP (UAS-P35 BL#5072, act>CD2>Gal4 from BL#4780)
w; tub>cd2>Gal4, UAS-CDB::GFP/CyO; tubGal80^{TS} (Buttitta et al., 2007)
y,w,hs-flp^12; + ; +
y,w,hs-flp^12; UAS-E2F, UAS-DP/Cyo-GFP; +
y,w,hs-flp^12; UAS-E2F, UAS-DP/Cyo-GFP; UAS-dMyc^{42}/TM6B (UAS E2F1,UAS DP from Neufeld et al 1998, dMyc from BL#9675)
y,w,hs-flp^12; UAS-E2F1, UAS-Dp/Cyo-GFP ; UAS-Tkv* (UAS Tkv* is UAS-Tkv^{Q235D} from (Nellen et al., 1996)
Pav::GFP (provided by Y. Yamashita) Ubiquitin-PAV::GFP; nanos-gal4

Fly rearing and mating:
All flies were raised and kept at room temperature (23°C) on Bloomington Cornmeal food unless otherwise noted. For experiments with virgins: males were collected as virgins and aged for indicated times in vials containing 7-10 males. For experiments with mated animals: males and females were kept at an approximate 1:1.5 ratio for indicated times.

Tissue fixation and staining:
Accessory glands were dissected in 1X PBS and were fixed in 4% PFA+1xPBS for 30 min at room temperature while rocking. Tissues were rinsed with 1xPBS+0.1%Triton-X twice for 10 minutes. Tissues were further permeabilized in 1xPBS+1.0%Triton-X for 30 minutes at room temperature while rocking. Tissues were rinsed with PAT for 10 min and primary antibodies diluted in fresh PAT were incubated at room temperature rocking overnight. Tissues were rinsed twice for 10 minutes in 1xPBS+0.1%Triton-X. Tissues were pre-blocked in PBT-X+2%NGS for 10 minutes. Secondary antibody was added to fresh PBT-X+2%NGS and tissues were incubated overnight rotating at room temperature. Tissues were rinsed twice in 1xPBS+0.1%Triton-X before incubating in DAPI (1µg/ml) for 10 min. Tissues were rinsed thoroughly before mounting with Vectashield.

The following antibodies were used in this study: Mouse anti DLG (DHSB) 1:500, Rabbit anti PH3 (Millipore #06-570) 1:1000, mouse anti PH3 (Cell Signaling #9706) 1:1000.

Measurements:
Fluorescent images were obtained using a Leica SP5 confocal, Leica SP8 confocal or Leica DMi6000B epifluorescence system. Brightfield images used to quantify gland size were taken on a Leitz Orthoplan.

To obtain overall gland size measurements, images were imported to Adobe Photoshop and Lasso Tool was used to outline the lobe of the accessory gland. Photoshop Measure was used to quantify pixels within the gland and measurements were transferred to Prism for analysis.

Discs large (DLG) antibody staining was used for all cell size measurements. Due to the apical localization of DLG, cell size measurements reported here are not of the volume of a cell, but rather a measurement of the apical area. Measurements reported here are taken mid-lobe and are only of main cells to ensure cell type differences are not confounding our measurements. Image J was used to obtain measurements of cells in microns and measurements were transferred to Prism for analysis.

Nuclear area measurements were done similarly to cell size (described above) using DAPI signal to delineate nuclei and include main cells only.

To measure levels of GFP, Cdt1-BFP and GFP-E2F1, Image J was used to measure integrated density of fluorescence within nuclei of secondary and main cells. Corrected total cell fluorescence (CTCF) was calculated following CTCF = Integrated Density – (Area of selected nuclei*mean fluorescence of background). [https://theolb.readthedocs.io/en/latest/imaging/measuring-cell-fluorescence-using-imagej.html](https://theolb.readthedocs.io/en/latest/imaging/measuring-cell-fluorescence-using-imagej.html)

**EdU labeling:**
Click-IT EdU AlexaFluor-555/488 Imaging kits were used as directed (Life Technologies). For labeling on the day of eclosion, accessory glands were dissected and immediately placed into Ringers solution containing 1mM EdU for 1 hour prior to fixing. For labeling during adult lifespan, animals were fed 1 mM EdU in 10% sucrose with blue food coloring for the indicated amounts of time. EdU/sucrose mixture was placed on whatman paper within empty vials and was changed every 2-3 days to control for contamination. We also performed feeding with 1 mM EdU in Cornmeal food with blue food coloring for up to 6 days and obtained similar results.

**DEMISE:**
Genotypes used:
y,w,hs-flp; DEMISE 10-3/+; Prd-Gal4/+  
y,w,hs-flp; DEMISE 10-3/+; +  
When kept at room temperature, on the day of eclosion animals already show DEMISE induced damage. For recovery experiments animals were collected as virgins and shifted to 18°C to reduce the Gal4/UAS expression levels. For EdU experiments animals were fed EdU/sucrose as described above for the indicated amounts of time.

Fertility assay: Individual virgin males were collected on the day of eclosion and were crossed with 15 Canton S virgin females (0-3 days of age). For DEMISE active: Virgin males were collected on the day of eclosion, crossed with virgin females, and kept at 25°C for the duration of the experiment. For DEMISE recovery: Virgin males were placed at 18°C for 5 day recovery period prior to mating and the flies were kept at 18 for the duration of the mating. For all experiments: Vials in which animals had laid in and were flipped out of were placed at 25°C to allow for proper development. Crosses were flipped every 2-3 days. Total number of progeny was counted.
Induction of growth regulators
Genotypes used:
y,w,hsflp; UAS-E2F1, UAS-DP/UAS-p35; act>CD2>gal4,UAS-GFP(nls)/+ 
y,w,hsflp; UAS-E2F1, UAS-DP/UAS-p35; act>CD2>gal4,UAS-GFP(nls)/ UAS-dMyc 
y,w,hsflp; UAS-E2F1, UAS-DP/ tub> Cd2>Gal4, UASCD8::GFP; UAS-Tkv*/UAS-tubGal80TS 
For EdU labeling: Newly eclosed flies were heat-shocked at 37°C for 60sec and aged for five days. Flies were fed EdU/sucrose as described above for 24 hours prior to dissection.

dMyc RNAi experiments
Genotypes used:
w:+; Prd-Gal4/UAS-dMyc-RNAi (BL25783/BL25784) 
w; UAS-dMYC-RNAi; Prd-Gal4 (BL51454) 
For Edu labeling: Both 1 hour incubation and 10 day feeding experiments were performed as described above. 
For nuclear area measurements: Measurements were performed as described above.

Sperm competition assays: Groups of 3-5 males were mated with W1118 females pre-mated with W1118. Males were placed with females within 24 hours of eclosion. Females were 1-5 days of age from bottles, all having mated previously (no virgin females were collected for this assay). Animals were left to mate for a total of 7 days and were flipped into new vials every 2-3 days. The percentages reported are the total number of orange-eyed offspring that emerge from each vial. All mating and development was done at 25°C.

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Figure 1: The adult accessory gland grows and main cells endoreplicate under normal physiological conditions throughout lifespan.

A: Adult accessory glands at Day 1 (day of eclosion) and Day 10 post eclosion. Accessory gland lobe is outlined with white dashed line. All glands measured are from virgins, so that size effects due to mating (i.e. release of seminal fluid proteins, hormonal signaling, muscle contractions, and emptying of the lumen) would not confound the measurements obtained.

B: Quantification of adult virgin male accessory gland area in pixels at indicated ages.

C: Quantification of adult virgin male accessory gland cell size and nuclear size in microns at indicated ages.

D: EdU incorporation assays in virgin male accessory glands. Accessory glands are outlined with yellow dashed line.

E: Magnification of virgin male accessory gland from 10 day EdU feeding at 20 day timepoint. EdU+ cells are main cells located near the distal tip of the gland. Secondary cells are outlined with yellow dashed line.

F: Magnification of mated male accessory gland. EdU+ cell is a main cell near the distal tip of the gland. Secondary cells are outlined with yellow.

Statistics are one-way ANOVA:

**** p < 0.0001
*** p = 0.0002

Scale bars:
A,D: 100 microns, E: 25 microns, F: 15 microns
**Figure 2: Cell cycle degradation machinery oscillates in the adult accessory gland.**


B: RGB cell cycle sensor in the adult accessory gland. Full length EGFP-PCNA to visualize early S Phase and cell cycle protein degrons nlsCycB1-96-nlsCycB1-285-tdTomato, to assay APC/C activity, and nlsCdt11-101-EBFP2, to assay CRL4 (Cdt2) activity. APC/C activity is high throughout the tissue, except in a small subset of cells, outlined in yellow. Few cells exhibit PCNA foci, indicating early S Phase, outlined in red. (Top panel) Mid lobe-region showing oscillations in main cells and (lower panel) distal tip showing oscillations in secondary cells.

C: Endogenous PCNA tagged with GFP in the adult accessory gland.

Scale bars:
A: 50 microns
B,C: 25 microns
**Figure 3: Main cells of the adult accessory gland are poised to endocycle.**

A: The Flippase-FRT system used to activate gene expression in clones in the adult accessory gland.

B: Quantification of adult accessory gland nuclear size with overexpression of E2F1/DP.

C: Overexpression of E2F/DP with dMYC and Tkv* in the adult accessory gland. White arrows: normal nuclear morphology, Yellow arrows: enlarged nuclei.

D: EdU incorporation with 1 day feeding in the adult accessory gland upon overexpression of dMyc at 4 days post eclosion.

Statistics are Welch’s T-test:
* p=0.0117

Scale Bars:
C,D: 25 microns
**Figure 4: dMyc is required for proper adult accessory gland endocycling and function**

A: Quantification of F1 progeny with orange eyes during sperm competition assay at 3 days, 5 days, and 7 days. Controls are crosses containing Prd-Gal4 without RNAi.

B: Adult accessory glands of 10 day virgin males. dMyc-i #1 (left) and dMyc-i #1 control. Control cross contains dMyc RNAi without a Gal4 driver.

C: Quantification of main cell nuclear size in 10 day virgin males dMyc-i #1, dMyc-i #1 control, dMyc-i #2, and dMyc-i #2 control. Control crosses contain dMyc RNAi without a Gal4 driver.

D: 1 hour EdU labeling on the day of eclosion in males expressing dMyc-i #1 using Prd-Gal4.

E: 10 day EdU/Sucrose feeding in virgin males expressing dMyc-i #1 using Prd-Gal4.

Statistics are one way ANOVA:

**** p<0.0001

Scale Bars:
B: 25 microns
D: 50 microns
Figure 5: The adult accessory gland can regenerate

A: Schematic of experimental procedure and representative outcomes using DEMISE system and a recovery protocol in the accessory gland. Red cells are representative of cells in which DEMISE is active and Reaper expression is high.

B: DEMISE system in the adult accessory gland at day of eclosion. Damaged lobe is outlined in yellow.

C: DEMISE system in the adult accessory gland at day 3 at room temperature. Damaged lobe is outlined in yellow.

D: DEMISE system in adult accessory gland that has been shifted to 18°C for 5 day recovery protocol.

E: DEMISE induced damage is visualized as pyknotic nuclei. Outlined with yellow dashed line is a binucleate accessory gland cell with two pyknotic nuclei.

F: Quantification of DEMISE system induced damage and accessory gland phenotype via categories of: normal size lobes, 1 damaged lobe, or 2 damaged lobes.

G: Adult accessory glands of 10 day mated males with the DEMISE system. Left two panels are animals mated at 25°C, right two panels are animals mated at 18°C after 5 day recovery at 18°C. Top panels are DEMISE control (DEMISE without a Gal4 driver), bottom panels are DEMISE driven by Prd-Gal4.

H: Total number of offspring for each mating condition: 25°C DEMISE active, 25°C DEMISE control, 18°C DEMISE recovery, and 18°C DEMISE recovery control. Controls are crosses containing DEMISE with no Gal4 driver.

Scale bars:
E: 25 microns
G: 100 microns
Figure 6: The adult accessory gland regeneration incorporates increased endoreplication and exhibits compensatory cellular hypertrophy

A: DEMISE induced damage with recovery protocol (Fig 5) has one normal lobe that has undergone very low levels of damage and one lobe that has undergone damage and responded with compensatory cellular hypertrophy (yellow arrow).

B: Magnification of both DEMISE accessory gland lobes from recovery protocol (Fig 5). One lobe with normal cell size (left) and one lobe that has undergone compensatory cellular hypertrophy (right).

C: Schematic of experimental procedure and representative outcomes using DEMISE system to induce damage and recovery protocol in the accessory gland paired with EdU feeding for different lengths of time.

D: DEMISE accessory gland from male that was fed EdU for the entire 11 day recovery protocol.

E: DEMISE accessory gland from male that was fed EdU for only the last 5 days of the 11 day recovery protocol.

F: Quantification of EdU+ main cells for a 7 day RT feeding and EdU feeding for the entire 11 day recovery protocol (approx. 5 days RT).

Statistics are Welch’s T Test:
** p=.0035

Scale Bars:
A: 50 microns
B: 25 microns
Figure 7: Developmental control of the cell cycle in the Drosophila accessory gland.

A: GFP negative cells show enlarged nuclei upon clone induction; these cells neighbor GFP-high cells.

B: Quantification of GFP levels from Fig 7A. Quantifications were taken of GFP-high cell, the GFP-low neighbor with the enlarged nuclei, and non-neighboring GFP negative cells (to subtract background). GFP in cells neighboring specific GFP-high cells is only visible when overexposing the GFP signal.

C: Pavarotti::GFP in the accessory gland. The localization of Pav::GFP (yellow arrow) suggests that main cells are in sister pairs from the penultimate cell cycle when the ring canal is formed. (Sister-pairs are shown here in a psuedocolored overlay.)

D: A model for variant cell cycle regulation in the developing accessory gland main cells. Progressive truncations of cytokinesis and mitosis culminate in endocycling that persists into adult stages.

Scale Bars:
C: 15 microns
Supplemental Figures

Supplemental Figure 1:

A: Quantification of fluorescence intensity of nlsCdt1^{1-102}-EBFP1 and GFP-E2F1_{1-230} from Figure 2A and B.

B: Alignment of the fly CDT1 N-terminal fragment in the RGB cell cycle reporter (Fig2) with the human CDT1 sequence.