Adipocyte amino acid sensing controls adult germline stem cell number via the amino acid response pathway and independently of Target of Rapamycin signaling in Drosophila

Alissa R. Armstrong1,2, Kaitlin M. Laws1,2 and Daniela Drummond-Barbosa1,2,3,*

ABSTRACT
How adipocytes contribute to the physiological control of stem cells is a critical question towards understanding the link between obesity and multiple diseases, including cancers. Previous studies have revealed that adult stem cells are influenced by whole-body physiology through multiple diet-dependent factors. For example, nutrient-dependent pathways acting within the Drosophila ovary control the number and proliferation of germline stem cells (GSCs). The potential role of nutrient sensing by adipocytes in modulating stem cells in other organs, however, remains largely unexplored. Here, we report that amino acid sensing by adult adipocytes specifically modulates the maintenance of GSCs through a Target of Rapamycin-independent mechanism. Instead, reduced amino acid levels and the consequent increase in uncoupled tRNAs trigger activation of the GCN2-dependent amino acid response pathway within adipocytes, causing increased rates of GSC loss. These studies reveal a new step in adipocyte-stem cell crosstalk.

KEY WORDS: Germline stem cells, Adipocytes, Amino acid transporters, Diet, Oogenesis, Drosophila

INTRODUCTION
Stem cell lineages are inextricably linked to whole-body physiology and nutrient availability in multiple organisms (Ables et al., 2012). For example, diet influences wound healing, hematopoietic transplants and cancer risk in humans, and evidence ranging from human epidemiological to model organism experimental data suggests that diet-dependent pathways impact a variety of adult stem cells (Ables et al., 2012). As intact living organisms vary their dietary input, multiple tissues and organs sense and respond to diet; however, our knowledge of how inter-organ communication contributes to the dietary control of adult stem cells remains limited.

The obesity epidemic has brought to light the crucial importance of normal adipocyte function in maintaining a healthy physiology. Adipocytes are highly sensitive to diet and produce long-range factors with key roles in metabolism, reproduction and other physiological processes (Rosen and Spiegelman, 2014). Conversely, dysfunctional adipocytes underlie the link between obesity and several diseases, including cancers (Vucenik and Stains, 2012). Whether sensing of dietary inputs by adipocytes leads to specific effects on adult stem cells in other organs, however, remains largely unexplored.

Drosophila female germline stem cells (GSCs) sense and respond to diet through complex endocrine mechanisms (Ables et al., 2012). Two or three GSCs reside within a well-defined niche in the germarium, the anterior region of the ovariole (Fig. 1A-C). Each asymmetric GSC division yields another GSC and a cystoblast that forms a 16-cell cyst, which is enveloped by follicle cells to generate a follicle that develops through oogenesis to form a mature oocyte (Spradling, 1993). On a yeast-rich diet, GSCs and their progeny grow and proliferate faster than on a yeast-free diet (Drummond-Barbosa and Spradling, 2001), and this response is mediated by diet-dependent factors that act on or within the ovary. For example, optimal levels of Target of Rapamycin (TOR) activity likely controlled by circulating amino acids are intrinsically required in GSCs for their proliferation and maintenance (LaFever et al., 2010; Sun et al., 2010). Insulin-like peptides produced by median neurosecretory cells in the brain act directly on GSCs to modulate how fast they proliferate to generate new cyts (LaFever and Drummond-Barbosa, 2005; Hsu et al., 2008). In parallel, insulin-like peptides act directly on cap cells, the major cellular components of the niche, to control GSC maintenance via two mechanisms. Insulin-like peptides promote the response of cap cells to Notch ligands (Hsu and Drummond-Barbosa, 2009, 2011), which are required for proper cap cell numbers (Song et al., 2007), and also GSC-cap cell attachment via E-cadherin (Hsu and Drummond-Barbosa, 2009, 2011). These past studies, however, did not address whether or how nutrient sensing by adipocytes influences the dietary response of GSCs and their descendants.

Drosophila adipocytes, together with hepatocyte-like oenocytes, compose the fat body (Fig. 1A), a nutrient-sensing organ with endocrine roles (Colombani et al., 2003; Arrese and Soulages, 2010; Rajan and Perrimon, 2012). In the larval fat body, TOR activation downstream of amino acid sensing results in the production of unknown factors that modulate overall growth of the organism (Colombani et al., 2003). In both the larval and adult fat body, sensing of sugars and lipids leads to the production of a leptin-like cytokine, Unpaired 2 (Upd2), which controls the secretion of brain insulin-like peptides (Rajan and Perrimon, 2012). Here, we report that partially inhibiting amino acid transport in adult adipocytes results in a specific reduction in the number of ovarian GSCs and that, surprisingly, this effect is independent of TOR signaling. Instead, reduced amino acid levels and the consequent increase in uncoupled tRNAs trigger activation of the GCN2-dependent amino acid response (AAR) pathway within adipocytes, causing increased rates of GSC loss. These results indicate that amino acid sensing by adipocytes through a TOR-independent mechanism is communicated to GSCs to control their maintenance, thereby contributing to their response to diet. Our
findings bring to light the importance of elucidating how adipocytes contribute to the regulation of various adult stem cell types by diet, and how these mechanisms might be adversely affected in obese individuals.

RESULTS

A tool for specific genetic manipulation of adult adipocytes

As a first step towards specific genetic manipulation of adult adipocytes using the UAS/Gal4/Gal80 system (del Valle Rodriguez et al., 2012), we sought to identify a Gal4 driver that, in adults, shows expression exclusively in adipocytes. We tested several Gal4 drivers with previously reported expression in the larval and/or adult fat body (Fischer et al., 1988; Colombani et al., 2003; Gronke et al., 2003; Rusten et al., 2004; Lazarova et al., 2007; DiAngelo et al., 2009) using a UAS-GFP reporter (green) driven by Gal80ts; Lsp2 shows robust expression in adipocytes on a rich diet at 29°C (D), but is not expressed either at 18°C (E) or on a poor diet (F). DAPI (blue) labels nuclei. Scale bar: 50 μm.

Decreased amino acid transport in adult adipocytes inhibits egg production

Amino acids are key dietary components that have systemic effects on organismal growth during development through their action in the larval fat body (Colombani et al., 2003). To test whether amino acid sensing in adipocytes might have an effect on the adult GSC lineage, we knocked down individual amino acid transporters in adult adipocytes using Gal80ts; Lsp2 and available UAS-RNAi lines (supplementary material Table S1, Fig. 2A). The Drosophila genome encodes 40 predicted amino acid transporters (www.flybase.org). Knockdown of single transporters in adult adipocytes did not lead to gross abnormalities in ovarian follicle development, fat body morphology or overall female health (Fig. 2B). For approximately one-third of the amino acid transporters tested, however, adipocyte-specific knockdown resulted in a significant decrease in the number of eggs laid (Fig. 3). These results suggest that incomplete loss of function of single transporters (and presumably relatively small changes in intracellular amino acid levels) within adipocytes are sufficient to influence oogenesis.

Reduced amino acid transport in adult adipocytes leads to increased GSC loss

Changes in GSC number or activity can contribute to alterations in egg production. We therefore determined whether GSCs are specifically affected by amino acid transport within adipocytes. Based on their pronounced egg laying reduction (Fig. 3) and reported fat body expression (supplementary material Table S1), we focused our analyses on females with adipocyte-specific knockdown of the amino acid transporters encoded by CG12773, slimfast (sif), CG7708, CG1607, CG1628 and CG13384. The number of GSCs declined significantly faster over time in females with adipocyte amino acid transporter knockdown relative to controls (Fig. 4A; supplementary material Fig. S3), suggesting that reductions in amino acid levels within adipocytes can be communicated to the ovary to influence GSC maintenance.

GSC loss downstream of adipocyte amino acid sensing is not a consequence of severe niche impairment or of alterations in systemic insulin signaling

We next tested whether decreased amino acid levels in adipocytes cause GSC loss through reduced bone morphogenetic protein (BMP) signaling from the niche, which is required for GSC maintenance (Xie and Spradling, 1998). We measured the nuclear levels of phosphorylated Mad (pMad), a reporter of BMP signaling (Kai and Spradling, 2003), and found that GSCs in adipocyte transporter knockdown females showed variable levels of pMad (Fig. 4B,C). Specifically, there is a small (less than 50%) decrease in pMad levels for three amino acid transporters (encoded by CG12773, CG13384 and CG1607), and an increase in those levels for the remaining transporters (CG1628, sif and CG7708). Even excluding sif and CG7708 (for which sample sizes are small) from this analysis, there is no consistent and drastic decrease in pMad levels, even though all six transporters share the same reduced
GSC number phenotype. It is therefore unlikely that a severe impairment of BMP signaling is the cause of GSC loss, in agreement with the observation that adipocyte knockdown of CG1628 shows a more severe GSC loss than knockdown of CG13384. We also did not observe dying cells adjacent to the GSC niche (Fig. 5A), suggesting that GSCs might be lost through differentiation, despite presumably adequate levels of BMP signaling.

During larval development, amino acid sensing by the fat body modulates systemic insulin signaling (Colombani et al., 2003), and our previous work showed that insulin-like peptides control GSC maintenance through the niche by controlling cap cell numbers and E-cadherin-mediated GSC-cap cell adhesion (Hsu and Drummond-Barbosa, 2009, 2011; Kwak et al., 2013). We therefore asked whether the reduction in GSC number resulting from amino acid transporter knockdown in adipocytes was due to changes in cap cell number or in E-cadherin levels. The numbers of cap cells, however, were unaltered (supplementary material Fig. S4A). Similarly, there were no obvious differences in the levels of E-cadherin at the niche-GSC junction (supplementary material Fig. S4B; number of germaria analyzed: 96 for control; 76 for CG1607; 15 for CG1628; 31 for CG7708; 74 for CG12773; 45 for CG13384; 69 for slif), although we cannot exclude the possibility of minor effects on E-cadherin levels based on this visual assessment. Also inconsistent with a general reduction in insulin signaling, GSC proliferation was increased and follicle cell division rates (a proxy for rates of follicle development) were unaltered or slightly increased in most cases upon amino acid transporter knockdown in adipocytes (Fig. 5B,C). Thus, reduced amino acid transport in adult adipocytes causes a specific decline in GSC numbers that is independent of changes in systemic insulin signaling or severe niche alterations.

Generally amino acid sensing in adult adipocytes does not affect vitellogenesis but appears to partially inhibit ovulation
To determine whether the GSC loss observed downstream of amino acid transporter knockdown in adipocytes is accompanied by additional alterations in the GSC lineage, we examined later stages of oogenesis. Onset of vitellogenesis and ovulation are major points of control of oogenesis by diet (Drummond-Barbosa and Spradling, 2001); therefore, we examined whether amino acid transport within adipocytes may also contribute to modulation of these processes. There was no increase in the percentage of ovarioles containing dying vitellogenic follicles upon adipocyte amino acid transporter knockdown, with the exception of slif antisense, which caused a small but significant increase in degeneration of vitellogenic follicles (supplementary material Fig. S5). These results are also consistent with normal levels of systemic insulin signaling, which are required for intact vitellogenesis (Drummond-Barbosa and Spradling, 2001; LaFever and Drummond-Barbosa, 2005; Hsu et al., 2008). By contrast, knockdown of several amino acid transporters resulted in a slight increase in the fraction of ovarioles showing a partial block in the ovulation of mature oocytes relative to controls (Fig. 6A,B). The partial block in ovulation, however, was a variable phenotype that did not reach statistical significance, presumably owing to the mild decrease in amino acid transport expected from knockdown of individual transporters.

Adipocyte TOR signaling controls ovulation but does not mediate the effects of adipocyte amino acid sensing on GSC maintenance
The nutrient sensor TOR acts downstream of Slif within larval adipocytes to promote organismal growth (Colombani et al., 2003), prompting us to ask whether adipocyte TOR signaling mediates the effects of amino acid transporters within adult adipocytes on the
ovarian GSC lineage. We inhibited TOR signaling specifically within adult adipocytes using \textit{Gal80^ts; Lsp2}-driven overexpression of the Tuberous Sclerosis Complex (Tsc)1/Tsc2 complex (Tapon et al., 2001) [a negative regulator of TOR (Laplante and Sabatini, 2012)] or of a dominant-negative version of RagA [\textit{RagAT16N} (Kim et al., 2008)], a positive regulator of TOR involved in amino acid sensing (Laplante and Sabatini, 2012). Inhibition of adipocyte TOR signaling using either of these established tools caused a marked increase in the percentage of ovaries showing a partial block in ovulation relative to controls (Fig. 6A,B). Adipocyte TOR inhibition, however, had no effect on GSC (or cap cell) numbers (Fig. 6C, supplementary material Fig. S6), suggesting that adipocyte amino acid levels control GSC maintenance independently of TOR signaling.

Increased levels of unloaded tRNAs and GCN2 activation in response to reduced amino acid levels in adipocytes cause GSC loss

We next hypothesized that the AAR pathway may act within adipocytes to control GSC numbers. The AAR pathway, conserved from yeast to mammals, senses limitations in one or more amino acids. Reduced amino acid levels lead to an increase in unloaded tRNAs, which activate the kinase GCN2, thereby controlling downstream translational and transcriptional events (Gietzen and Rogers, 2006; Gallinetti et al., 2013; Bjordal et al., 2014). Our hypothesis therefore predicts that directly increasing unloaded tRNA levels in adipocytes should reduce GSC numbers. Inhibiting or mutating aminoacyl-tRNA synthetases (the enzymes responsible for coupling amino acids to their cognate tRNAs) are well-established approaches to experimentally increase uncharged tRNA levels, thereby activating the AAR pathway under normal amino acid levels (Gietzen and Rogers, 2006; Gallinetti et al., 2013; Bjordal et al., 2014). Given that Slif, a cationic amino acid transporter (Colombani et al., 2003), is among those that function in adipocytes to control GSC numbers (see Fig. 4A), we knocked down the genes encoding Histidyl-, Arginyl- or Lysyl-tRNA synthetases (\textit{Aats-his}, \textit{Aats-arg} or \textit{Aats-lys}, respectively) in adipocytes using \textit{Gal80^ts; Lsp2} (supplementary material Fig. S7). Control or \textit{Aats-lys} adipocyte knockdown (which was relatively inefficient; supplementary material Fig. S7C) had no effect on GSC number. By contrast, knockdown of aminoacyl-tRNA synthetases using either an
Aats-his or two distinct Aats-arg RNAi transgenes in adipocytes led to a marked decrease in GSC number upon transgene induction (Fig. 7A, supplementary material Fig. S7A). Similar to what we observed for amino acid transporter knockdown, there were no changes in cap cell number (supplementary material Fig. S7B). These results show that activation of the AAR pathway suffices to phenocopy the GSC loss caused by reduced amino acid transport in adipocytes. Conversely, adipocyte-specific knockdown of Gcn2 reverts the GSC loss caused by RNAi of the amino acid transporter CG12773 (Fig. 7B,C), further suggesting that the AAR pathway is also required to mediate the effects of adipocyte amino acid transporters on GSCs. Based on these results, we conclude that the

Fig. 4. Adult adipocyte-specific knockdown of amino acid transporters leads to increased rates of GSC loss in the ovary. (A) Average number of GSCs per germarium at 0, 5 or 10 days of Gal80ts; Lsp2-mediated induction of RNAi or antisense transgenes against amino acid transporters or white control. See supplementary material Fig. S3 for sample sizes and distribution. **P<0.01; ****P<0.0001, two-way ANOVA with interaction. Data shown as mean±s.e.m. (B) Germaria at 10 days of adipocyte-specific GFP control or amino acid transporter RNAi labeled for phosphorylated Mad (pMad; green), 1B1 (red, fusome) and Lamin C (red, cap cell nuclear envelope). GSC nuclei are outlined. Scale bar: 2.5 μm. (C) Box and whisker plot of mean nuclear pMad intensity for experiment in B. Sample sizes are included above. **P<0.01; ***P<0.001, Student’s t-test.

Fig. 5. Reduced amino acid transport in adipocytes does not affect cell death within the germarium, but causes a slight increase in GSC proliferation. (A) Germaria from females at 10 days of adult adipocyte-specific knockdown of CG1628 or white control showing some occurrence of cell death (ApopTag, green) in both cases. DAPI (blue) labels nuclei; 1B1 (red) labels fusomes; LamC (red) labels cap cell nuclear envelopes. Scale bar: 10 μm. In the graph on right, bars represent the percentage of germaria containing ApopTag-positive cells, with the hatched region indicating the fraction of those displaying ApopTag adjacent to GSC niche. The number of germaria analyzed is shown above each bar. (B,C) Frequencies of GSCs (B) or follicle cells (C) in S phase, based on EdU incorporation, at 10 days of adipocyte knockdown of amino acid transporters or GFP control. Number of GSCs (B) or follicle cell fields (C) analyzed is shown above each bar. *P<0.05; **P<0.01; ***P<0.001, Student’s t-test. Data shown as mean±s.e.m.
AAR pathway within adipocytes is sufficient and required to initiate an amino acid-dependent signaling cascade of inter-organ communication to modulate GSC maintenance in the ovary.

DISCUSSION

The specific effects of adipocyte dysfunction on normal stem cell lineages have remained largely unexplored. Yet, clear evidence shows that obesity leads to higher risk for multiple chronic diseases (Vucenik and Stains, 2012). Our data support the model that amino acid levels within adipocytes are sensed through separate mechanisms that specifically affect a stem cell lineage at distinct stages (Fig. 7D). The AAR pathway acting within adipocytes influences maintenance of GSCs, whereas amino acid sensing through the adipocyte Rag/TOR pathway modulates the efficiency of ovulation of fully differentiated GSC daughters, or oocytes. Future studies should identify the extracellular factors acting downstream of these intra-adipocyte signaling cascades to communicate adipocyte nutritional status to the GSC lineage. This work underscores the importance of investigating the role of inter-organ communication in the control of stem cells and their differentiated daughters in a wide variety of systems. Furthermore, it suggests that the aberrant co-option of endocrine pathways that normally tie stem cell lineages to whole-body physiology might contribute to the increased cancer risk associated with obesity (Vucenik and Stains, 2012).

Drosophila as a model for investigating how inter-organ communication contributes to the regulation of adult stem cells

Drosophila is an ideal model for molecular physiology studies, owing to the ease of cell/tissue-specific manipulations (del Valle Rodriguez et al., 2012), which are essential to dissect how individual systemic signaling events contribute to complex physiological networks. Indeed, recent years have seen an explosion in metabolism and physiology studies using Drosophila (Colombani et al., 2003, 2012; Gutierrez et al., 2007; Arquier et al., 2008; Geminard et al., 2009; Palanker et al., 2009; Sieber and Thummel, 2009; Slaidina et al., 2009; Delanoue et al., 2010; Ruaud et al., 2011). Particularly useful throughout these studies is the UAS/Gal4 system, which allows tissue- and/or cell-type-specific genetic manipulations; however, a crucial consideration when designing such studies is the specificity of Gal4 expression to avoid misinterpretation of phenotypes. Indeed, most of the published fat body drivers we tested were not expressed exclusively in adipocytes in adult females. By contrast, the robust and highly specific expression of 3.1Lsp2-Gal4 in adipocytes makes it a valuable tool for exclusive genetic manipulation of adipocytes to test how they impact not only GSCs, but also other adult stem cell types.

In addition to adipocytes, nutrient sensing by other tissues also affects GSCs. For example, insulin-like peptides secreted from the brain act directly on the germline to modulate GSC proliferation, cyst growth and vitellogenesis, and also indirectly affect GSC maintenance through effects on the niche (LaFever and Drummond-Barbosa, 2005; Hsu and Drummond-Barbosa, 2009, 2011). Other adult stem cell types are also modulated by insulin signaling, including male GSCs and intestinal stem cells (Ables et al., 2012). Much remains unknown, however, about how other tissues influence stem cells, despite evidence suggesting endocrine roles for muscle (O’Brien et al., 2011; Demoniti et al., 2014), intestines (Reiher et al., 2011) and the brain (Nassel and Winther, 2010).

Separate modes of amino acid sensing in adipocytes affect the stem cell lineage at distinct stages

Our findings that amino acid sensing by adipocytes controls GSC maintenance through the AAR pathway and ovulation through TOR clearly illustrate the high degree of specificity of adipocyte-to-ovary communication. Our results also imply that relatively small fluctuations in amino acid levels (e.g. those resulting from partial knockdown of single amino acid transporters) within adipocytes can be effectively transmitted to the ovary to modulate stem cell number. These same slight reductions in amino acid levels resulted in less significant effects on ovulation, consistent with the distinct amino acid sensing mechanisms involved. It will be very interesting to identify and study the effectors downstream of AAR and TOR signaling that mediate these distinct effects on the GSC lineage.

Not surprisingly, inhibition of TOR signaling impacted ovulation more severely than manipulation of single amino acid transporters, in agreement with its role downstream of transporters and as an integrator of multiple inputs, including nutrients, energy status and growth factors (Dibble and Manning, 2013). It is likely that additional stimuli upstream of TOR within adipocytes also regulate ovulation.

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**Fig. 6. GSC loss induced by adult adipocyte-specific knockdown of amino acid transporters is independent of TOR signaling.** (A) Ovaries at 10 days of adipocyte-specific CG12773 knockdown or Tsc1/Tsc2 induction showing retention of mature oocytes in subsets of ovarioles (‘blocked’ ovulation). Mature oocytes are recognizable by the presence of dorsal appendages (arrowheads). Scale bar: 500 μm. (B) Percentage of ovaries containing at least one ovariole that retains more than one mature oocyte at 10 days of adipocyte-specific amino acid transporter knockdown (left) or at different days of inhibition of TOR signaling (right). **P<0.01. Data from 0 d and 15 d time points are from one experiment, whereas 10 d represents three experiments. Number of ovaries analyzed is shown above each bar. (C) Average number of GSCs at different days of Gal80D; Lsp2-mediated induction of a dominant-negative RagA (RagAT16N) or of Tsc1/Tsc2 transgenes showing that inhibition of TOR signaling has no effect on GSC maintenance. See supplementary material Fig. S6 for sample sizes and distribution. Control is GFP RNAi for A and B (left), and Gal80D; Lsp2 alone for B (right) and C.
The AAR pathway is evolutionarily conserved from yeast to humans; however, its downstream targets are context dependent. In yeast, for example, phosphorylation of eIF2α by activated GCN2 causes selective upregulation of translation of the transcriptional factor GCN4, which in turn induces genes involved in amino acid transport as well as in amino acid biosynthesis (Natarajan et al., 2001). Translational derepression of ATF4 (the GCN4 equivalent in Drosophila and humans), by contrast, leads to expression of oxidative stress genes in mouse embryonic fibroblasts (Harding et al., 2003). The targets of the AAR pathway in the context of intact multicellular organisms remain largely unidentified. Nevertheless, it is reasonable to speculate that the sets of targets regulated by the AAR pathway in different tissues and cell types may be quite different, given the diversity of processes being modulated. For example, the AAR pathway acts in the brains of Drosophila larvae, mice and rats to reduce intake of food sources that lack essential amino acids (Hao et al., 2003). The targets of the AAR pathway in the context of intact organisms contribute to the control of GSC maintenance.

**Context-specific targets of the amino acid response pathway**

Adipocytes, stem cells and increased cancer risk in obese individuals

Obesity and high calorie intake are associated with increased risk of multiple cancer types, including breast, colon and prostate cancer (Bianchini et al., 2002; McMillan et al., 2006; Xue and Michels, 2010). Similar to GSCs and other stem cells (Ables et al., 2012), cancers are highly responsive to nutrient-sensing pathways, and components of the insulin and TOR pathways are often misregulated in cancers (Jee et al., 2005; Chen, 2011). Given the parallels between cancer cells and stem cells, investigations of the role of adipocytes in adult stem cell regulation will likely provide valuable insights into the link between obesity and cancer risk. Based on our results, we speculate that aberrant communication of the nutrient-sensing status of fat cells could modulate the activity of cancer cells and might explain the link between diet, adiposity and cancer.

**MATERIALS AND METHODS**

**Drosophila strains and culture conditions**

Fly stocks were maintained at 22-25°C on standard medium containing cornmeal, molasses, yeast and agar. Standard medium supplemented with wet yeast paste was used for all experiments, except for Fig. 1F, where flies were kept on molasses/agar plates with no yeast. Previously described fat body Gal4 lines were used: Adh-Gal4 (Fischer et al., 1988), cg-Gal4 (Rusten et al., 2004), FB-Gal4 (Gronke et al., 2003), r4-Gal4 (DiAngelo et al., 2009), pumpless-Gal4 (Colombani et al., 2003) and 3.1Lsp2-Gal4 (Lazareva et al., 2007). The temperature-sensitive tub-Gal80⁰ transgene has been described (McGuire et al., 2003). UAS-RNAi lines obtained from the
Vienna Drosophila RNAi Stock Center (http://stockcenter.vdrc.at) and the Transgenic RNAi Project (http://www.flyrna.org) collection at Bloomington Drosophila Stock Center (http://flystocks.bio.indiana.edu) for knockdown of amino acid transporters are listed in supplementary material Table S1. Other UAS-RNAi lines used were: P(GD14098)v42184 (line 1) and P(GD14098)v42185 (line 2), against Aats-arg; P(KK102374), VIE-260B, against Aats-his; P(TRIP.HMS00763)attP2, against Aats-lys; P(TRIPT.GL00267)antP2, against Gcn2; P(UAS-GFP,dxRNAi.R)143, against GFP (used as a control); P(TRIP.JF01545)antP2, and against white (used as a control). The UAS-CG17273(G4); UAS-Gcn2(Rbh); and tub-Gal80(gal80); 3.1Lsp2-Gal4 (Gal80p; Lsp2) lines were generated by standard crosses. The following UAS lines have been described previously: UAS-silf(antense (Colombani et al., 2003), UAS-Tsc1; UAS-Tsc2 (Tapon et al., 2001) and UAS-Rag(F10) (Kim et al., 2008). Other genetic elements used are described in FlyBase (http://www.flybase.org).

Adult adipocyte-specific genetic manipulations

For adult adipocyte-specific genetic manipulation, females of genotypes yw; tub-Gal80+/+; 3.1Lsp2-Gal4/UAS-X or yw; tub-Gal80(UAS-X; 3.1Lsp2-Gal4/+ were used. (UAS-X represents any of the UAS transgenes in this study.) Females were raised at 18°C, the permissive temperature for Gal80p, to keep transgene expression off during development. Newly eclosed females were maintained at 18°C for 3 days and then switched to 29°C, the restrictive temperature for Gal80p, for various lengths of time to induce transgene expression prior to dissection and/or analyses.

Immunostaining and fluorescence microscopy

All tissues were dissected in Grace’s medium (BioWhittaker) and fixed in 5.3% formaldehyde (Ted Pella) in Grace’s medium at room temperature for the following amounts of time: 13 min for ovaries, 20 min for abdominal carcasses (containing attached fat body) or brains, and 1 h for guts. Tissues were rinsed and washed three times in 0.1% Triton X-100 (Sigma) in phosphate-buffered saline (PBS), or PBT, and subsequently blocked in 5% bovine serum albumin (BSA; Sigma) and 5% normal goat serum (NGS; Jackson ImmunoResearch) in PBT, or blocking solution, for 3 h at room temperature or overnight at 4°C. Tissues were incubated overnight at 4°C in the following primary antibodies diluted in blocking solution: rabbit anti-GFP (Torrey Pines; 1:2500, TP401); mouse monoclonal anti-Hits (1B1) (DSHB; 1:10); mouse anti-α-spectrin (3A9) (DSHB; 1:50); mouse monoclonal anti-Lamin C (LC28.26) (DSHB; 1:100); rat monoclonal anti-E-cadherin (DCA2D) (DSHB; 1:100); and rabbit anti-pMad (Smad3) (Epitomics; 1:100, #1880). [This particular Smad3 antibody is widely used in Drosophila to detect pMad specifically (Hayashi et al., 2009; Ables and Drummond-Barbosa, 2010; Issigonis and Matunis, 2012; Matsuoka et al., 2013; Ma et al., 2014; Sulkowski et al., 2014.)] Tissues were washed in PBT, and incubated for 2 h at room temperature in 1:200 Alexa Fluor 488- or 568-conjugated secondary antibodies (Molecular Probes). Samples were washed, and ovaries, brains, guts and fat bodies (scraped off from carcasses) were mounted in Vectashield containing DAPI (Vector Labs). For visualization of lipid droplets, fixed and blocked carcasses were incubated in 1:200 Nile Red (Sigma) in 50% glycerol in PBS for 10 min at room temperature. Fat bodies were mounted in 90% glycerol in PBS containing 0.5 μg/ml DAPI (Sigma). Data were collected with a Zeiss Axiosimager-A2 fluorescence microscope or a Zeiss LSM700 confocal microscope. For nuclear pMad quantification, the densitometric mean of individual GSC nuclei was measured from optical sections containing the largest nuclear diameter (visualized by DAPI) using AxioVision. (To achieve as much consistency as possible among samples for pMad measurements, ovaries were dissected, fixed and stained in parallel under identical conditions, and the image acquisition settings were exactly the same for all images used for quantification.)

GSC and cap cell analyses

Cap cells were identified based on their ovoid shape and Lamin C staining, and GSCs were identified based on their juxtaposition to cap cells and fusome morphology and position, as described previously (Hsu et al., 2008; Hsu and Drummond-Barbosa, 2009). For statistical analysis of differences in rates of GSC loss we used two-way ANOVA with interaction (www.graphpad.com), which, simply stated, calculates the significance of any differences measured among genotypes in how much GSC numbers change over time.

Egg counts and ovulation analyses

To measure egg production, five pairs of flies (females of appropriate genotype and y w wild-type males) were maintained in plastic bottles containing molasses/agar plates covered by a thin layer of wet yeast paste, in triplicate, at 29°C. Plates were replaced daily, and eggs laid within the preceding 24 h were counted on specific days throughout experiments.

For ovulation analyses, females were dissected in Grace’s medium and intact ovaries were examined under a Zeiss Stemi 2000 stereomicroscope. Each ovariole in a wild-type ovary typically contains zero or one mature oocyte, recognizable by its fully developed dorsal appendage (Spradling, 1993). Ovaries in which at least one ovariole contained two or more mature oocytes were classified as having partially blocked ovulation. Images of whole ovaries were captured using a Nikon Coolpix L620 digital camera.

EdU incorporation, apoptosis assay and quantification of vitellogenesis defects

For EdU analysis, intact ovaries were incubated for 1 h at room temperature in 100 μM EdU (Molecular Probes) diluted in Grace’s medium, washed, fixed as described and permeabilized for 20 min in 0.5% Triton X-100 in PBS. Following primary antibody incubation, EdU samples were subjected to the Click-IT reaction according to the manufacturer’s protocol (Life Technologies) for 30 min at room temperature. GSC proliferation rates were determined by calculating the fraction of EdU-positive GSCs as a percentage of the total number of GSCs analyzed per genotype. To measure follicle cell proliferation, single confocal planes transecting follicle monolayers (i.e. follicle cell fields) at the top and bottom of flatly mounted ovarioles were acquired, and the average percentage of EdU-positive follicle cells per follicle cell field was calculated. This analysis included follicle cells covering follicle stages 4 to 6, prior to the mitotic-to-endoreplication switch, as described previously (LaFever et al., 2010).

ApopTag Direct In Situ Apoptosis Detection Kit (Millipore) was used as described (Drummond-Barbosa and Spradling, 2001). Progression through vitellogenesis was assessed using DAPI staining (Spradling, 1993). Ovarioles containing vitellogenic follicles were easily distinguished from those with blocked vitellogenesis, which contained at least one dying vitellogenic follicle. Dying vitellogenic follicles were recognizable by their position within the ovariole and by the presence of pyknotic nuclei.

RT-PCR analyses

Fat bodies from 2-10 females per genotype at 10 days of RNAi induction were hand dissected in RNALater solution (Ambion). RNA was extracted using the RNAqueous-4PCR DNA-free RNA Isolation for RT-PCR kit (Ambion) and cDNA was synthesized using the SSRII kit (Ambion) according to the manufacturer’s protocols. For each primer pair, PCR was performed on both the control and corresponding RNAi samples. The primers used are listed in supplementary material Table S2. Rp49 primers were used as a control. Band intensity was quantified using AxioVision by background pixels from band pixels in a fixed size box (net band intensity) and normalized to the net band intensity of the corresponding Rp49 band. Controls were set to one and experimental sample intensities were determined relative to control.

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Competing interests

The authors declare no competing financial interests.
Author contributions A.R.A. and D.D.-B. designed the experiments and wrote the manuscript. A.R.A. and K.M.L. performed the experiments. A.R.A., K.M.L. and D.D.-B. interpreted the experiments.

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