

#### **RESEARCH ARTICLE**

# Drosophila Mon1 constitutes a novel node in the brain-gonad axis that is essential for female germline maturation

Neena Dhiman<sup>1,2</sup>, Kumari Shweta<sup>1</sup>, Shweta Tendulkar<sup>2</sup>, Girish Deshpande<sup>3</sup>, Girish S. Ratnaparkhi<sup>2,\*</sup> and Anuradha Ratnaparkhi<sup>1,\*</sup>

#### **ABSTRACT**

Monensin-sensitive 1 (Mon1) is an endocytic regulator that participates in the conversion of Rab5-positive early endosomes to Rab7-positive late endosomes. In Drosophila, loss of mon1 leads to sterility as the mon1 mutant females have extremely small ovaries with complete absence of late stage egg chambers - a phenotype reminiscent of mutations in the insulin pathway genes. Here, we show that expression of many Drosophila insulin-like peptides (ILPs) is reduced in mon1 mutants and feeding mon1 adults an insulin-rich diet can rescue the ovarian defects. Surprisingly, however, mon1 functions in the tyramine/ octopaminergic neurons (OPNs) and not in the ovaries or the insulinproducing cells (IPCs). Consistently, knockdown of mon1 in only the OPNs is sufficient to mimic the ovarian phenotype, while expression of the gene in the OPNs alone can 'rescue' the mutant defect. Last, we have identified ilp3 and ilp5 as critical targets of mon1. This study thus identifies mon1 as a novel molecular player in the brain-gonad axis and underscores the significance of inter-organ systemic communication during development.

KEY WORDS: *Drosophila* oogenesis, Insulin production, Inter-organ communication, Neuronal control

#### INTRODUCTION

Drosophila oogenesis has served as an attractive and genetically tractable developmental model system with numerous distinguishing features (Matova and Cooley, 2001; McLaughlin and Bratu, 2015; Roth and Lynch, 2009; Spradling et al., 2001). The two prominent traits that make it simple yet unique include exquisitely detailed patterning orchestrated by the dialogue between the germline and the surrounding soma, and also fine-tuned coordination between the non-autonomous and autonomous factors that determine morphogenesis and growth (Li and Xie, 2005; Lin and Spradling, 1993; Roth and Lynch, 2009).

Drosophila adult ovary is made up of a pair of bundles, each containing 16-20 tubular structures termed ovarioles (Margolis and Spradling, 1995; Robinson and Cooley, 1997; Roth and Lynch, 2009). Each ovariole is an independent egg assembly line, comprising of six to eight egg chambers that develop in a sequential manner and are connected by stalk cells. The growth of the egg chamber is broadly divided into the previtellogenic (i.e. up to stage 8) and vitellogenic

<sup>1</sup>Agarkar Research Institute (ARI), Pune, India. <sup>2</sup>Indian Institute of Science Education & Research (IISER), Pune, India. <sup>3</sup>Department of Molecular Biology, Princeton University, Princeton, NJ 08540, USA.

 $\verb|^*Authors| for correspondence (girish@iiserpune.ac.in; anu.aripune@gmail.com)|$ 

K.S., 0000-0002-5692-905X; G.S.R., 0000-0001-7615-3140; A.R., 0000-0001-6548-6185

In this regard, we turned our attention to a highly conserved endocytic protein, Monensin sensitive 1 (hereafter referred to as Mon1). A protein complex between Mon1, a 'longin domain'-containing protein, and CCZ1 functions as a guanine nucleotide exchange factor

(GEF) for Rab7 (Nordmann et al., 2010). In a canonical endocytic cycle, Mon1 appears to be essential for the conversion of Rab5-positive early endosomes to Rab7-positive late endosomes (Nordmann et al.,

phases (i.e. beyond stage 8). The vitellogenic phase is distinguished by the exponential growth of the oocyte due to yolk accumulation. The transition from pre-vitellogenic to vitellogenic stage is under tight hormonal, as well as nutrient, control. Juvenile hormone (JH) is required for yolk formation and is necessary for vitellogenesis (Gilbert et al., 1998; Mirth et al., 2014; Raushenbach et al., 2004; Wilson, 1982); the nutritional status of the fly and the dependent growth control is regulated by insulin-like peptides (Grönke et al., 2010; Richard et al., 2005; Mirth et al., 2014; Mendes and Mirth, 2016), which also play an important role in this process. An additional layer of complexity is brought about by interactions between ecdysone-JH and insulin signaling in regulating this crucial check-point (Gruntenko and Rauschenbach, 2008, 2018; Ikeya et al., 2002; Shim et al., 2013).

Drosophila encodes eight insulin-like peptides (ILPs), of which three members (ILP2, ILP3 and ILP5) are produced by the median neurosecretory cells (MNCs) or insulin-producing cells (IPCs) (Ikeya et al., 2002). These insulin-like peptides are released into circulation through the axon terminals of the IPCs at the corpora cardiaca – a neurohemal organ (Brogiolo et al., 2001; Cao and Brown, 2001; Ikeya et al., 2002). Supporting the conclusion that ILPs exert a non-autonomous influence on the progression of oogenesis, germline-specific loss of Insulin receptor (INR) or Chico (the insulin substrate protein) leads to a reduction in the size of the ovary, an inability of the egg chambers to enter the vitellogenic stages and, consequently, sterility (Drummond-Barbosa and Spradling, 2001; LaFever and Drummond-Barbosa, 2005; Richard et al., 2005).

Previous studies in *Drosophila* have indicated that IPCs are likely to be under neuronal regulation: short neuropeptide F (sNPF) and octopamine appear to stimulate IPCs and thus can potentiate insulin signaling, whereas GABA has an inhibitory influence (Enell et al., 2010; Lee et al., 2008; Luo et al., 2014). Despite the fact that insulin signaling affects a wide variety of developmental processes in several organismal contexts, mechanisms underlying insulin production, release and transport are not fully understood. The modes of short- as well as long-range transmission have been a focus of enquiry owing to the systemic influence of insulin signaling on growth and metabolism (Erion and Sehgal, 2013; Kenyon, 2010; Partridge et al., 2011). It is generally believed that different regulatory circuits, both upstream and downstream, of insulin signaling must deploy a unique and dedicated set of regulators to achieve tissue-specific outcomes. The molecular circuitry, which participates during the synthesis and long-distance transmission of insulin, to engineer proper Drosophila egg chamber growth and patterning has yet to be elucidated.

2010; Poteryaev et al., 2010). Consequently, loss of Mon1 leads to accumulation and enlargement of early Rab5-positive endocytic compartment. In *Drosophila*, the process of Rab conversion is conserved and *mon1* is involved in the recruitment of Rab7 (Yousefian et al., 2013). Intriguingly, at the neuromuscular junction neuronal *mon1* is involved in regulating glutamate receptor levels on the post-synaptic side (Deivasigamani et al., 2015).

While characterizing the P-element mediated excision mutation in *mon1*, we noticed that homozygous mutant animals die throughout development whereas the escapers are short lived with severe motor defects (Deivasigamani et al., 2015). Recently, we also observed that *mon1* mutants display sex-non-specific sterility. Expectedly, the lethality and motor defects could be rescued by pan-neuronal expression of *mon1*. Curiously, however, the neuronal expression was also capable of rescuing the sterility in these mutants, suggesting that the neuronal expression of *mon1* may regulate fertility in wild-type females.

Here, we have analyzed mechanistic underpinnings of the influence of *mon1* on oogenesis. Our findings have uncovered an unanticipated link between neuronal *mon1* activity in octopaminergic/

tyraminergic neurons (OPNs) and insulin production/signaling from the IPCs that has functional implications for the brain-gonad axis.

#### **RESULTS**

#### mon1 mutations influence body size and ovarian size

We have previously reported the generation of new alleles of *mon1* using P-element excision (Deivasigamani et al., 2015). Molecular characterization of one of these alleles,  $mon1^{\Delta 181}$ , revealed that the C-terminal region from amino acid 249 is deleted. Homozygous  $mon1^{\Delta 181}$  mutants die throughout development and display motor defects (Deivasigamani et al., 2015). In addition,  $mon1^{\Delta 181}$  mutant adults are ~15% smaller in size (Fig. 1A,B) compared with wild type and exhibit sex non-specific sterility. On average, wild-type females are are 2535.82±72.27 µm (n=10) in length, whereas  $mon1^{\Delta 181}$  mutant females measure 2188±97.07 µm (n=10) (Fig. 1A,B). Sterility is fully penetrant with homozygous  $mon1^{\Delta 181}$  females laying very few eggs, if any.

As a first step towards understanding the underlying cause for sterility, we dissected ovaries from 2- to 3-day-old  $mon1^{\Delta I8I}$  mutant females. The mutant ovaries were remarkably smaller than those in

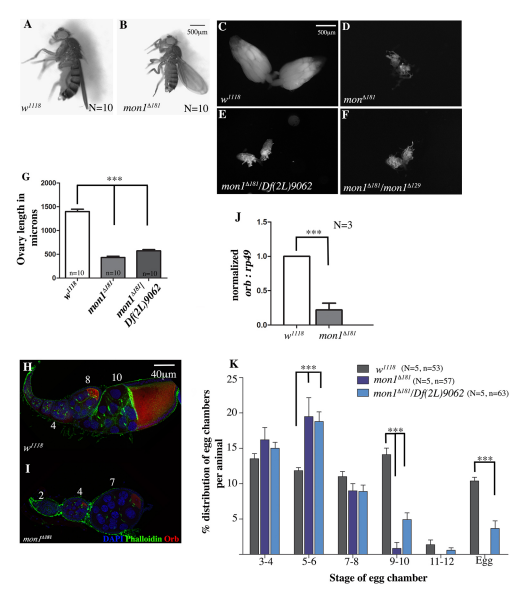


Fig. 1. mon1 mutants have smaller ovaries with egg chambers that are stalled at pre-vitellogenic stages. (A,B) Two- to 3-day-old wild-type (A) and age-matched  $mon1^{\Delta 181}$  animals. The mutant is smaller, reduced by ~15% in length, on average, when compared with wild type. (C-F) Ovaries from 2- to 3-day-old virgin females of wild type (C), homozygous  $mon1^{\Delta 181}$  (D),  $mon^{\Delta 181}$ / Df(2L)9062 (E) and  $mon^{\Delta 181}/mon^{\Delta 129}$ (F). Mutant ovaries are extremely small compared with wild type. (G) Measurement of ovary length along the proximal-distal axis. Mutants show a ~70% reduction in size. (H,I) Wild-type (H) and homozygous mon<sup>∆181</sup> (I) ovarioles stained for Orb (red), and with DAPI (blue) and phalloidin (green). Late-stage egg chambers (stage 9-14) tend to be absent in mon1 mutants (J). Orb transcript levels are reduced by ~80%, with late stage egg chambers (stage 9-14) tending to be absent in  $mon 1^{\Delta 181}$  mutants. (K) Bar graph showing the distribution of egg chamber types from stage 3 onwards in 2- to 3-day-old  $w^{1118}$ , homozygous  $mon^{4181}$ and  $mon^{\Delta 181}/Df(2L)9062$  adult virgin females. Early pre-vitellogenic egg chambers predominate in the mutants. N=number of animals. n=number of ovarioles; \*\*\*P<0.001, two-way ANOVA.

control animals (Fig. 1D). To exclude the possibility that this phenotype is genetic background dependent, we also examined ovary size in  $mon1^{\Delta I8I}/Df(2L)9062$  and  $mon1^{\Delta I8I}/mon1^{\Delta I29}$  females.  $mon1^{\Delta I29}$  carries a small deletion that spans the C-terminal region of mon1 and the 5' end of the neighboring smog gene (Deivasigamani et al., 2015). In both cases, the ovaries were found to be very small and comparable in size with  $mon1^{\Delta I8I}$  mutants (Fig. 1E,F).

To estimate the extent of decrease in ovary size, we measured the length of the wild-type and mutant ovaries from the proximal to distal end and found that  $mon1^{A181}$  ovaries (431.02±27.6  $\mu$ m; n=10) were approximately one-third of the size of the wild-type control (1400±49.9  $\mu$ m, n=10; Fig. 1G). A comparable decrease in size was also observed in ovaries from  $mon1^{A181}/Df(2L)9062$  adults (571.5±26  $\mu$ m, n=10; Fig. 1G). Furthermore, the total number of ovarioles per animal was also significantly reduced: compared with 35.05±0.4 (n=10) in wild type, the average number of ovarioles in  $mon1^{A181}$  mutants was reduced by ~35% (21.3±1.13; n=5).

### mon1 mutant ovaries exhibit arrested egg chamber development and partial degeneration

To assess whether different ovarian cell types (somatic as well as germline) are correctly specified, ovaries from *mon1* females were stained with DAPI, a DNA dye and fluorescently tagged phalloidin to mark actin-rich structures. We also co-immunostained these samples with antibodies against an oocyte marker, Orb. Ovaries from age-matched wild-type animals were used as controls.

The most striking phenotype seen in these mutants was the absence of late stage egg chambers (Fig. 1I). Most mutant egg chambers appeared to be arrested at pre-vitellogenic stage, with the terminal egg chamber being of stage 7 or 8 (Fig. 1I). Ten percent of the mutant ovarioles also showed degenerating terminal egg chambers (data not shown). Orb levels were also reduced compared with the control (Fig. 1H,I). Consistent with a reduction in protein levels, orb mRNA levels were diminished by 80% in  $mon1^{\Delta 181}$  ovaries (normalized transcript abundance of 1 in wild-type versus  $0.22\pm0.1$  in  $mon1^{\Delta 181}$  mutants; Fig. 1J).

The developmental arrest was quantified by plotting the percentage distribution of the egg chamber stages across ovarioles (Fig. 1K). Egg chambers counted were clubbed for two consecutive stages, e.g. stages 3 and 4 or 5 and 6 for clearer representation. In contrast to wild type (Fig. 1K),  $mon1^{\Delta 181}$  mutants, show fewer egg chambers stage 8 onwards with a near absence of egg chambers beyond stage 9 (Fig. 1K). A similar distribution of egg chambers was observed in the ovaries from  $mon1^{\Delta 181}/Df(2L)9062$  with a very low proportion of ovarioles displaying late stage egg chambers and mature egg formation (Fig. 1K). Taken together, these data suggest that loss of mon1 leads to a developmental arrest during oogenesis, resulting in the stalling of egg chamber growth at stage 7-8.

### Ovary-specific knockdown of Mon1 is insufficient to recapitulate the ovarian phenotype of *mon1* mutants

For a better understanding of the developmental arrest in *mon1* mutant ovarioles, we sought to identify the cells or tissues responsible for the phenotype by reducing *mon1* transcripts in the ovary using RNAi. *mon1* expression was reduced in the germline and somatic epithelial sheath cells by using *nanos* (*nos*)-*Gal4* and *heartless(htl)* <sup>GMR93H07</sup>-*Gal4* (Irizarry and Stathopoulos, 2015) drivers, respectively. In both cases, the ovaries appeared comparable in size to their respective controls, and ovarioles with mature eggs could be seen (Fig. 2A-D). The distribution of egg chambers stages in ovaries from RNAi and control animals was similar (Fig. 2E,F) suggesting that reduction of *mon1* in the ovary did not affect egg

chamber development. Other ovary-specific drivers such as *E22C-Gal4* (which is expressed in both the follicle cells and germline) and maternal *alpha-tubulin-*GAL4:*VP16* (*mat-Gal4*) were also used for knockdowns and in both instances we failed to observe an effect on ovary size and egg chamber growth (Fig. S1A,B), leading us to conclude that ovarian tissue is unlikely to be the site of Mon1 action.

#### Downregulation of *mon1* in octopaminergic neurons leads to a decrease in ovary size and delays maturation of egg chambers

We have previously reported that pan-neuronal expression of *mon1* is able to rescue lethality as well as the neuromuscular junction phenotype induced by loss of *mon1* (Deivasigamani et al., 2015). We therefore wondered whether neuronal downregulation of *mon1* could recapitulate some aspects of the ovarian phenotype of *mon1*<sup>Δ181</sup>. To test this, we expressed *UAS-mon1RNAi* in the nervous system using the pan-neuronal driver *C155-GAL4*. Surprisingly, this resulted in a significant reduction in ovary size (Fig. 2H), which was evident from the substantial increase in aspect ratio of these ovaries (compare Fig. S1C,D with Fig. S1E). We examined the distribution of egg chambers in these ovaries and found a near 50% decrease in ovarioles with mature eggs (Fig. 2K), which partially recapitulates the ovary phenotype seen in the mutants.

The ovary directly receives innervation at the ovarian peritoneal sheath muscles from the OPNs (Kurz et al., 2017; Middleton et al., 2006; Rezával et al., 2014). Knockdown of mon1 using tdc2-GAL4, which expresses in OPNs (Burke et al., 2012; Erion et al., 2012), resulted in smaller ovaries (Fig. 2I,J) with an average reduction of 22% in aspect ratio compared with control ovaries (Fig. S1F). A comparison of the distribution of egg chambers in tdc2-GAL4>mon1RNAi ovaries showed a significant increase in the proportion of pre-vitellogenic egg chambers along with a near 64% reduction in mature eggs (Fig. 2L). Thus, knockdown of mon1 in a subset of neurons (-the OPNs) seemed sufficient to partially mimic important aspects of the mon1 mutant phenotype. This suggests that *mon1* function in OPNs is required for escaping developmental arrest. The absence of a stronger and a more complete penetrance of the mutant phenotype is likely due to inefficient knockdown of mon1 transcript.

## Expression of mon1 in the OPNs is sufficient to rescue the ovarian defects of $mon1^{\Delta181}$ mutants

These data indicate that Mon1 activity in neurons, more specifically in the OPNs, influences egg chamber maturation. We therefore examined whether the ovary phenotype was rescued in mutant animals that expressed *UAS-mon1:HA* in all neurons given that pan-neuronal expression of the transgene rescues lethality (Deivasigamani et al., 2015). Indeed, the ovaries from the rescued animals appeared normal in size (compare Fig. 3B and Fig. 1C) and displayed late stage egg chambers and mature eggs.

Next, we tested whether expression of mon1, in the OPNs alone, using tdc2-GAL4 could rescue the lethality and sterility defect in  $mon1^{\Delta 181}$ . Surprisingly, a complete rescue of lethality was observed. Although homozygous mutant flies fail to survive beyond 7 days, the 'rescued' females were able to live well beyond 30 days (Fig. 3E). Moreover, the morphology of the ovaries in these animals was found to be remarkably similar to wild type (Fig. 3D) and the ovariole number was also restored in these animals (35.10±0.43, n=10,  $w^{1118}$ ) versus 25.30±0.84 (n=10,  $mon1^{\Delta 181}$ , tdc2-GAL4/ $mon1^{\Delta 181}$ ) versus 34.60±0.43 (n=10,  $mon1^{\Delta 181}$ , tdc2-GAL4/ $mon1^{\Delta 181}$ ; UAS-mon1-HA) (Fig. 3F).

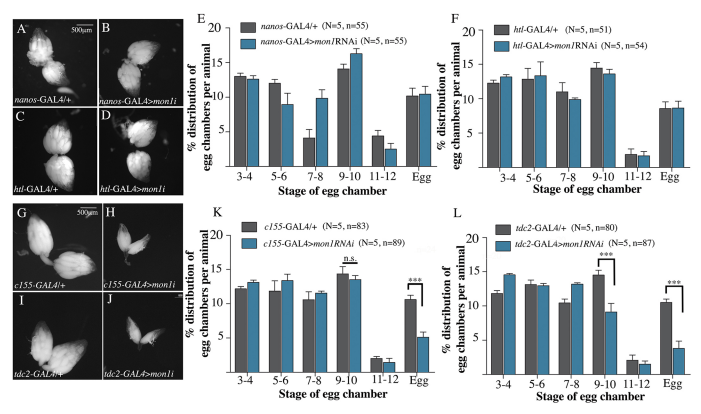


Fig. 2. Knockdown of *mon1* in OPN neurons partially phenocopies the developmental stalling seen in *mon1*<sup>4181</sup> ovarioles. (A-D) Downregulation of *mon1* in the germline (*nos*-GAL4>*UAS-mon1RNAi*) or sheath cells surrounding the ovary (*htl*<sup>GMR93H07</sup>-GAL4>*UAS-mon1RNAi*) does not reduce ovary size or perturb gross ovarian morphology. (E,F) Distribution of egg chambers in 2- to 3-day-old *nanos*-GAL4>*UAS-mon1RNAi* (E) and *htl*<sup>GMR93H07</sup>-GAL4> *UAS-mon1RNAi* (F) females is comparable with their respective controls. (G-J) Pan-neuronal (H) and *tdc2*-GAL4-specific (J) knockdown of *mon1* results in smaller ovaries compared with their respective controls (G,I). (K,L) Distribution of egg chamber types in 2- to 3-day-old *C155*-GAL4>*UAS-mon1RNAi* (K) and *tdc2*-GAL4>*UAS-mon1RNAi* (L) females along with their respective controls. Pan-neuronal knockdown of *mon1* leads to a 50% decrease in the number of mature eggs (10.61±0.62% in control versus 5.11±0.76% in RNAi animals). Expression of *UAS-mon1RNAi* in the *tdc-2* domain leads to a 37% decrease in stage 9-10 egg chambers (14.51±0.71% in control versus 9.12±1.26% in RNAi animals) and a 63% decrease in the number of eggs (10.51±0.5% in control versus 3.8±1.07% in RNAi animals). *N*=number of animals, *n*=number of ovarioles; \*\*\**P*<0.001, two-way ANOVA.

We quantified the distribution of egg chambers in these animals (tdc2-GAL4, mon1<sup>Δ181/Δ181</sup>>UAS-mon1-HA) and found a complete suppression of the developmental arrest: whereas in mutant females about 3% of the egg chambers are in stage 9-10, in 'rescued' females this frequency was 14% (Fig. 3G), which is comparable with wild-type control females (compare purple bar in Fig. 3G with gray bar in Fig. 1K). The ovaries of these females also showed the presence of mature eggs at a frequency of 10%, which is similar to that seen in wild-type females. A more detailed examination of the ovarioles by staining them with anti-Orb, phalloidin and DAPI showed presence of late stage egg chambers, eggs, the complete absence of degenerating of egg chambers normally seen in the mutants and the restoration of orb expression.

To determine the extent to which the observed rescue might be due to 'leaky' expression of the UAS transgene, we also analyzed the percentage distribution of egg chambers in  $mon1^{\Delta 181/\Delta 181} > UAS$ -mon1:HA animals. A minor increase in the percentage of egg chambers of stage 9-10 was observed (2.9% in  $mon1^{\Delta 181/181}$ , tdc2-GAL4 versus 6.82% in  $mon1^{\Delta 181/\Delta 181} > UAS$ -mon1:HA). These animals also showed the presence of eggs at a low frequency of 2%. However, in both cases the change did not appear to be statistically significant, indicating that 'leaky' expression is not a major contributor to the suppression of the ovary defect seen upon reconstitution of mon1 expression in the OPNs.

We further examined the fecundity of the 'rescued' females with age-matched wild-type females by crossing each of them individually to wild-type ( $w^{II18}$ ) males. Eggs laid by a single female were counted from the second day post-eclosion (Fig. 3H). Wild-type males were crossed to virgin females on the day of eclosion. Mutant  $mon1^{\Delta 181}$  females, when mated to wild-type males, do not lay eggs. In contrast, a single wild-type female lays between 50 and 65 eggs on the second day. From the 3rd day, this number increases to an average of ~85-95 eggs. Interestingly, the number of eggs laid by 'OPN rescue' females was comparable with wild type, with no significant difference in the number of eggs laid (Fig. 3H). In contrast, mutant females rescued using C155-GAL4 produced relatively fewer eggs, with the number of eggs being almost constant in the range of 65-70 after the second day. However, in both cases, as seen with wild-type females, more than 85% of the eggs were fertilized evident from the development of denticle belts. In addition, very few of these embryos hatched into first instar larvae, indicating a possible maternal requirement for mon1 during embryogenesis. Taken together, these results suggest that expression of mon1 in a small subset of  $\sim$ 137 neurons (the OPN neurons; Pauls et al., 2018)) alone is sufficient to rescue lethality and the ovary defect in  $mon1^{\Delta 181}$ , enabling oogenesis to progress beyond the previtellogenic stages, revealing an unanticipated, yet close, functional connection between Mon1 activity and the OPNs.

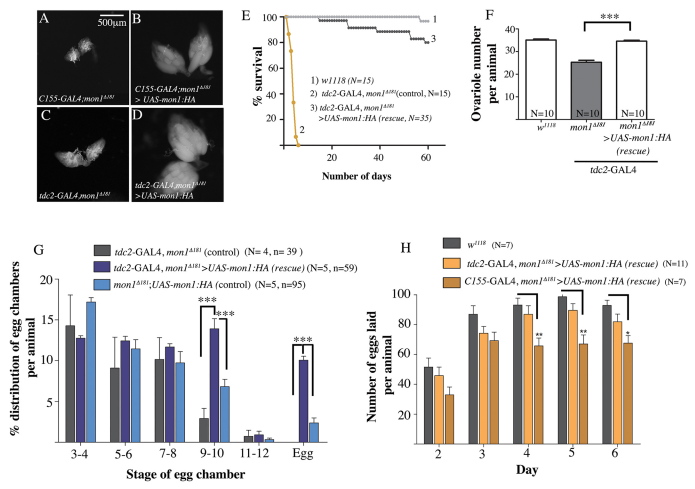


Fig. 3. Expression of mon1 in the OPNs is sufficient to rescue lethality and ovarian defects in  $mon1^{\Delta 181}$ . (A,B) Pan-neuronal expression of UAS-mon1:HA using C155-GAL4 in  $mon1^{\Delta 181}$  (B) rescues the 'small ovary' phenotype (A). (C,D) Expression of UAS-mon1:HA in a small subset of ~137 neurons, using tdc2-GAL4 (D, tdc2-GAL4,  $mon1^{\Delta 181}$ / $mon1^{\Delta 181}$ ; UAS-mon1:HA) also rescues the ovary phenotype in  $mon1^{\Delta 181}$ (C). (E) The short lifespan of  $mon1^{\Delta 181}$  adults is rescued by expression of the gene in the tdc2 domain (tdc2-GAL4,  $mon1^{\Delta 181}$ >; UAS-mon1:HA). The animals live for more than 30 days compared with the mutants that fail to survive beyond 10 days. (F) Expression of mon1:HA in OPNs also rescues the decrease in ovariole number observed in  $mon1^{\Delta 181}$  animals ( $35\pm0.433$  in wild type versus  $21.3\pm1.12$  in  $mon1^{\Delta 181}$  versus  $34.6\pm0.4$  in tdc2-GAL4,  $mon1^{\Delta 181}$ >UAS-mon1:HA). (G) The 'stalling' defect in  $mon1^{\Delta 181}$  ovarioles is rescued by expression of mon1:HA in the OPNs. More than a 4-fold increase is seen in the percentage of stage 9-10 egg chambers with respect to the 'GAL4' control ( $2.91\pm1.23\%$  in mutant 'GAL4' control versus  $33.9\pm1.24\%$  in 'rescue') and a 2-fold increase with respect to the 'UAS' control ( $6.8\pm0.9\%$  in 'UAS' mutant control versus  $33.9\pm1.24\%$  in  $33.9\pm1.24\%$  i

### Expression of *Drosophila* insulin-like peptides is perturbed in *mon1* mutants

The role of insulin signaling during ovarian development in *Drosophila melanogaster* is well established (LaFever and Drummond-Barbosa, 2005) and arrested oogenesis is a hallmark of insulin pathway mutants. For example, mutations in *chico*, the substrate for insulin receptor, also cause sterility and prevent transition of the egg chambers from pre-vitellogenic to the vitellogenic phase (Richard et al., 2005) resulting in smaller ovaries.

Given the broad similarity in the ovarian phenotypes between  $mon1^{\Delta 181}$  and insulin pathway mutants, we wondered whether mon1 activity might target one or more of insulin-like peptides. We therefore decided to test whether endogenous insulin levels and/or signaling are compromised in mon1 mutants. There are eight insulin-like peptides (ILPs) in Drosophila. As a first test,

we quantitated the expression levels of these ILPs in total mRNA samples made from mutant female adults using quantitative RT-PCR. While no significant change was observed in the expression of *ilp2* and *ilp7*, a significant decrease was noted in the case of *ilp1*, *ilp3*, *ilp4*, *ilp5* and *ilp6*. Curiously, a strong increase in expression was noted for *ilp8* (Fig. 4A).

Given the significant reduction in ILP gene levels in mon1 mutants, we wondered whether feeding the adult mutants an insulinrich diet would ameliorate the oogenesis defects. One of the striking effects seen upon transferring newly eclosed  $mon1^{\Delta 181}$  mutant females to medium containing 5  $\mu$ g/ml of insulin (see Materials and Methods) was the improvement in their lifespan. While most homozygous  $mon1^{\Delta 181}$  mutants die within 5 to 7 days of eclosion, mutants maintained on insulin-containing medium were able to survive for 15 days post-eclosion (Fig. 4B). We examined ovaries of

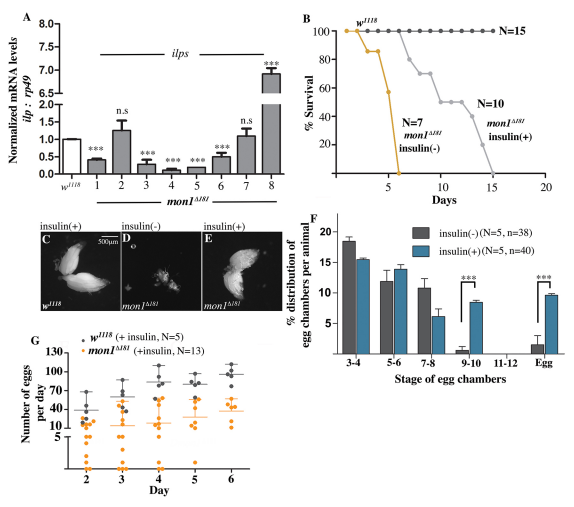


Fig. 4. Feeding insulin to  $mon1^{\Delta181}$  suppresses ovary defects. (A) Ilp mRNA levels from 2- to 3-day-old adult  $mon1^{\Delta181}$  whole animals normalized to wild type using quantitative real-time PCR. ilp1, ilp3, ilp4, ilp5, ilp6 and ilp8 show significant changes in transcript levels in the mutants. (B) Lifespan of  $mon1^{\Delta181}$  maintained on normal  $[mon1^{\Delta181}$  insulin(-)] and 5  $\mu$ g/ml insulin-containing  $[mon1^{\Delta181}$  insulin(+)] medium. The half-life of survival is nearly doubled in animals fed on insulin-containing medium. (C-E) Five-day-old ovaries from wild type (C, insulin positive) and  $mon1^{\Delta181}$  flies (D,E) kept on normal (D) and insulin-containing (E) medium. Ovary size is restored in mutants fed on medium supplemented with insulin (E). (F) Percentage distribution of egg chamber stages in  $mon1^{\Delta181}$  [insulin(-)] and  $mon1^{\Delta181}$  [insulin(+)] animals. There is a significant increase in the number of stage 9-10 egg chambers [insulin(-): 0.59±0.59 versus 8.47±0.32 in insulin(+) animals] and in the number of eggs [insulin (-): 1.49±1.49 versus 9.6±0.27 in insulin(+) animals]. (G) Comparison of the number of eggs laid by  $mon1^{\Delta181}$  flies fed with insulin. Each dot represents eggs laid by a single animal.  $mon1^{\Delta181}$  flies usually do not lay eggs. However, when kept on sucrose media supplemented with insulin, the females start laying  $mon1^{\Delta181}$  give should be a single animal.  $mon1^{\Delta181}$  flies usually do not lay eggs. However, when kept on sucrose media supplemented with insulin, the females start laying  $mon1^{\Delta181}$  gives half the number achieved by wild-type females.  $mon1^{\Delta181}$  flies is a supplemented with insulin, the females start laying  $mon1^{\Delta181}$  gives half the number achieved by wild-type females.  $mon1^{\Delta181}$  flies is a supplemented with insulin, the females start laying  $mon1^{\Delta181}$  gives half the number of ovarioles.

5-day-old insulin-fed mutants with age-matched mutants maintained on normal medium. In contrast to mutant ovaries (Fig. 4D), the ovaries from the insulin-fed flies appeared to be nearly normal (Fig. 4E) in size and comparable with ovaries from wild-type animals fed on insulin (Fig. 4C). Importantly, ovarioles from insulin-fed flies displayed late-stage egg chambers (Fig. 4F). The average percentage of egg chambers at stage 9-10 in insulin-fed flies was ~8.5% compared with 0.6% in control mutants. Similarly, the percentage of mature eggs increased dramatically to 10%, as seen in wild-type animals (see Fig. 2F), compared with 1.5% in 'unfed' controls.

To confirm that the effects of insulin are not due to an increase in the amount of protein in the medium, we carried out experiments in which mutant flies were separately maintained either on yeast-rich medium or on medium containing an equivalent amount of FLAG peptide (Sigma, F3290). In both cases, the ovaries appeared similar to those from mutant animals maintained on normal medium. This suggests that the suppression of the ovary phenotype in insulin-fed

flies is not due to a nutrient or amino-acid rich diet but likely because of insulin activity.

To determine the physiological relevance of insulin feeding, we sought to test whether fecundity in the mutants was restored by assaying individual mutant females for their ability to lay eggs upon insulin feeding. Wild-type flies fed with insulin were used as controls. Whereas all insulin-fed  $mon1^{\Delta 181}$  flies were able to lay eggs in their lifetime, there was considerable variation in the number of eggs laid by individual females, presumably due to variability in feeding. Moreover, because the assays were conducted on insulincontaining sucrose agar medium, 60% of the mutant females failed to survive beyond the third day. Nonetheless, about 20% of the flies were able to produce, on an average, more than 35 eggs per day, with some females laying as many as 40 eggs (Fig. 4G). Furthermore, 80% of the eggs laid appeared to be fertilized, based on development of cuticular structures. The observation that feeding insulin is able to alleviate both the morphological aberrations and the functional defects suggests that the effect of insulin on ovarian

growth is not simply a result of an 'insulin-dependent bypass'. These data thus argue that the insulin pathway is likely a physiologically relevant target of *mon1* activity.

#### ILP5 levels are considerably reduced in mon1 mutant IPCs

The transcript levels of a number of ILPs are reduced in *mon1* mutants (Fig. 4A). Taken together with the observation that an insulin-rich diet can partially suppress the ovarian phenotype, we wondered whether *mon1* influences ILP levels. We therefore examined whether *mon1* is specifically necessary to maintain ILP levels in the IPCs. IPCs synthesize ILP2, ILP3 and ILP5 but only *ilp3* and *ilp5* mRNA levels are reduced in *mon1* mutants. As *ilp5* has been shown to influence oogenesis (Badisco et al., 2013; Richard et al., 2005), we decided to examine the level of ILP5 in IPCs by staining the adult brains using anti-ILP5 antibodies. Compared with wild

type, a substantial decrease in ILP5 levels was observed in  $mon1^{\Delta 181}$  IPCs and in the axons projecting from these cells (compare Fig. 5A with 5B,C). In some animals, the decrease was very dramatic, with IPCs displaying ILP5 staining only at the periphery (Fig. 5C) of the cell body. We quantified the intensity of ILP5 staining in the IPCs and compared it with the levels in the heterozygous mutant and wild-type animals. In both cases, a 50-60% decrease in ILP5 intensity was observed. Given the association between Mon1 and the OPNs in the context of ovary maturation (Fig. 2), we wondered whether knockdown of mon1 in the OPNs might affect ILP5 levels. Interestingly, expression of mon1 RNAi in OPNs led to a significant decrease in ILP5 staining in the IPCs as well as the projecting axons (compare Fig. 5E with 5F). Curiously, the staining in the RNAi animals appeared punctate. A quantification of the staining intensity showed a 40% decrease in RNAi animals compared

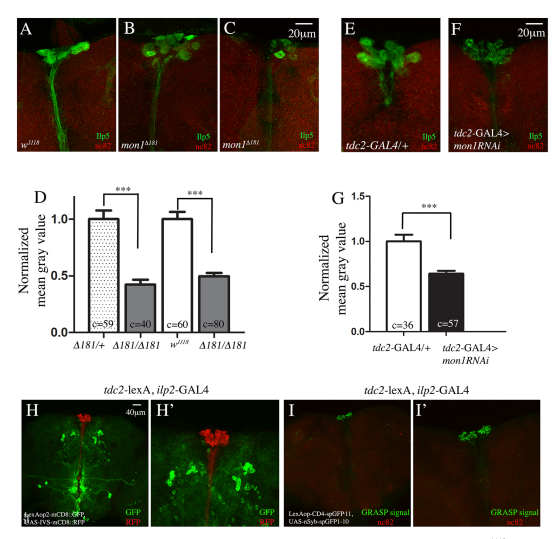


Fig. 5. mon1 in OPNs regulates ilp5, and OPNs make direct synaptic connections with the IPCs. (A-C) Adult brains of  $w^{1118}$  (A) and  $mon1^{\Delta181}$  mutants (B,C) stained using anti-ILP5 (green) and anti-Brp (red) antibodies. There is a distinct decrease in the intensity of ILP5 staining in the mutants (B,C). In a few cases, the IPCs exhibit only faint staining around the edge of the cells (C). (D) Quantification of the staining intensity shows, on average, a 50% reduction in staining intensity between control and mutants  $[mon1^{\Delta181}]$ , 1.0±0.07 (c=59 cells, N=6) versus 0.43±0.04 (c=40 cells, N=6) in  $mon1^{\Delta181}$ ;  $w^{1118}$ , 1.0±0.06 (c=60 cells, N=6) versus 0.49±0.3 (c=80 cells, N=7) in  $mon1^{\Delta181}$ ]. (E-G) Adult brains of tdc2-GAL4/+ (E) and tdc2-GAL4>VAS-mon1RNAi (F) stained using anti-ILP5 (green) and anti-Brp (red) antibodies. Intensity of ILP5 staining in the IPCs is reduced in upon downregulation of mon1 in OPNs (compare F with E). (G) The decrease in intensity is ~40% [tdc2-GAL4/+, 1.00±0.07 (t=36, t=36) versus 0.63±0.03 (t=36) cells, t=360 in t=361 in t=362-GAL4>UAS-mon1RNAi]. (H,H') Adult brain of a t=362-lexA, t=363 in t=363 in t=363 and anti-GFP (green) and anti-GFP (red) antibodies. The organization of the two sets of neurons is shown (H). A more detailed image of H is shown in H'. (I,I') Adult brain of a t=363 in t=364 staining representing the GRASP signal is seen at the IPCs alone (I). A more detailed image of I is shown in I'.

with controls (Fig. 5G). Thus, both *ilp5* mRNA and protein levels are lowered in *mon1* mutants and knockdown of *mon1* in OPNs alone is sufficient to decrease the level of ILP5 in the IPCs.

Based on the proximity of the dendrites of the IPCs and the OPNs, it has been speculated that these neurons may form direct synaptic connections with each other (Luo et al., 2014). Given the non-autonomous regulation of *ilp5* by Mon1, we decided to test whether IPCs and OPNs form synaptic connections using the 'GFP resonstitution across synaptic partners' (GRASP) technique in which two membrane-bound complementary split fragments of GFP are expressed independently in two neuronal cell types to reconstitute a functional GFP at synaptic sites (Feinberg et al., 2008). To achieve this, we recombined the tdc2-LexA driver with *ilp2*-GAL4. We confirmed the presence of both drivers by crossing the recombinant line with GFP and RFP reporters for GAL4 and LexA, respectively (Fig. 5H,H'). Interestingly, when crossed to a line carrying the split-GFP reporters for LexA and GAL4, reconstituted GFP or GRASP signal was detected only at the IPCs (Fig. 5I,I'), indicating that the OPNs make direct synaptic contacts with the IPC cell body. This observation strongly suggests that the regulation of *ilp5* by Mon1 in the OPNs is likely to be direct.

### Overexpression of *ilp5* or *ilp3* in IPCs can alleviate loss of *mon1*

Based on the above results, we sought to assess whether the reduction in ILP5 levels in mon1 mutant IPCs is an important factor that contributes to its ability to regulate egg chamber maturation. If so, we expected neuronal expression of ILP5 to mitigate the ovarian phenotype due to loss of mon1. Consistent with this, expression of UAS-ilp5 using a pan-neuronal driver (C155-Gal4; compare Fig. 6A,B) or an IPC-specific driver (ilp2-Gal4; compare Fig. 6C with D) was able to ameliorate the ovary phenotypes of  $mon1^{\Delta 181}$ . With both drivers, nearly 50% of the mutant animals showed near wild-type-like ovaries, while others showed a partial rescue in which ovaries were bigger than  $mon1^{\Delta 181}$  mutants but with relatively fewer eggs. Furthermore, overexpression of ilp5 in the IPCs was able to restore ovariole number in the mutants ( $31.67\pm1.2$ , n=6). Improved viability was evident from the increase in frequency of obtaining mutant flies.

We compared the distribution of egg chambers between individual animals expressing *UAS-ilp5* and *UAS-mon1HA* under the pan-neuronal *C155*-GAL4. As expected, the latter showed substantial increase in the number of late-stage egg chambers and eggs. Expression of *ilp5* also led to an increase in the percentage of late-stage egg chambers and eggs, albeit to a lesser extent compared with *UAS-Mon1HA* (Fig. 6E).

In addition to ILP5, IPCs are known to synthesize ILP2 and ILP3. Previous studies have indicated existence of partial redundancy among the different ILPs (Broughton et al., 2008; Grönke et al., 2010). We examined whether pan-neuronal or IPC-specific overexpression of ilp2 and ilp3 could achieve an analogous rescue of the ovary phenotype in  $mon1^{\Delta 181}$ . Overexpression of ilp2 did not yield viable adults. In contrast, overexpression of ilp3 using C155-GAL4 suppressed the ovary defect in a manner comparable with ilp5 in terms of the ovariole number as well as relieving the egg chamber arrest (Fig. 6E). Similar to expression of ilp5, improvement in viability was evident from the increase in frequency of getting mutant adults.

Further supporting these results, overexpression of *ilp5* and *ilp3* in the IPCs alone was sufficient to suppress the egg chamber arrest. As with pan-neuronal expression, the overall ovarian morphology in 50% of the animals resembled wild type whereas in the others, the rescue was partial with an appreciable increase in the number of

late-stage egg chambers and eggs (Fig. 6F). Overexpression of *ilp5* and *ilp3* in the OPNs was unable to suppress the loss of viability and ovarian defects in the mutants, thus confirming the IPC-specific role of *ilp5* and *ilp3* (data not shown).

To determine whether expression of *ilp3* and *ilp5* also rescued fecundity, we assayed the ability of mutant females to lay eggs by individually crossing them to wild-type males. Indeed, these females were able to lay eggs and more than 75% of these eggs were fertilized (Fig. S2). An observation made during these egg-laying assays was the increase in mortality of mutant flies when placed on sucrose agar medium when compared with regular corn-meal agar medium, indicating increased sensitivity to nutrition. In summary, neuronal expression of both *ilp5* and *ilp3* in *mon1* mutant females was able to relieve the egg-chamber arrest during oogenesis, leading to restoration of the normal physiological function of the ovary. The partial nature of the rescue suggests presence of additional Mon1 targets or the requirement for combinatorial ILP expression. Future studies will focus on identification and characterization of these novel molecular players.

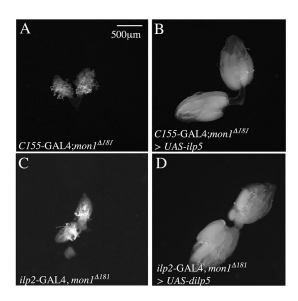
#### **DISCUSSION**

ILPs, in conjunction with the insulin receptor are involved in a variety of conserved biological processes that ultimately impact growth advantage, developmental profile, reproductive potential and aging-dependent decline (Boulan et al., 2015; Richard et al., 2005; Shim et al., 2013). Recent data also suggest that insulin can affect behavioral traits via nervous system function and physiology (Graham and Pick, 2017; LaFever and Drummond-Barbosa, 2005).

In Drosophila, differential expression of the ILPs and the systemic nature of their function underlie their diverse spatiotemporal roles (Bai et al., 2012; Grönke et al., 2010; Gruntenko and Rauschenbach, 2018; Hsu and Drummond-Barbosa, 2009; Knoblich, 2012; Richard et al., 2005; Shim et al., 2013; Ueishi et al., 2009). Emphasizing the importance of the long-distance nature of its communication and regulation, elegant studies by Rajan and Perrimon have demonstrated that Unpaired2 (Upd2) secreted by the fat body communicates with the IPCs to induce secretion of ILPs for maintenance of metabolic homeostasis (Rajan and Perrimon, 2012). A subsequent study from the Leopold lab has shown GPCR signaling by Methuselah (Mth) in response to Stunted (Sun) regulates secretion of ILPs (Delanoue et al., 2016). Regulation of insulin signaling during organ development and maturation has been studied extensively in the context of oogenesis and its role in regulating vitellogenesis is well established. In this study, we show that Mon1, an endocytic factor, regulates ovary maturation by participating in a non-autonomous mechanism to control expression of ILPs in the IPCs.

*mon1* mutants display female sterility; the mutant females have small ovaries with fewer ovarioles. The individual mutant egg chambers are arrested at the previtellogenic stages, resulting in ovaries that are devoid of any mature eggs. Interestingly, *mon1* activity appears to be required not in the ovaries, but in the nervous system. Supporting such a non-cell autonomous control by Mon1, compromising *mon1* levels specifically in the OPNs appears sufficient to induce developmental stalling at the vitellogenic checkpoint.

The observation that expression of *mon1* using a pan-neuronal driver is sufficient to rescue both lethality and sterility strongly supports a neural basis for the *mon1*-dependent defects. That expression in the approximately 176 OPNs labeled by *tdc2*-GAL4 in the brain and ventral ganglion (Pauls et al., 2018) is sufficient to rescue both lethality and sterility in these mutants argues that these might be the functionally crucial subset in this regard.



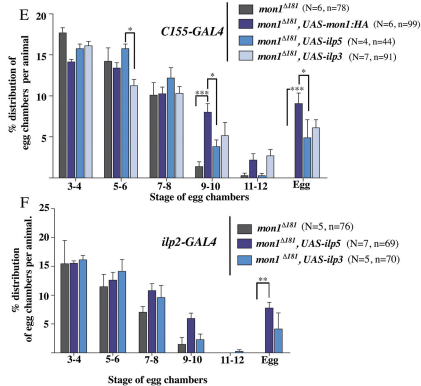


Fig. 6. Expression of ilp5 and ilp3 rescues the egg-chamber stalling defect in  $mon1^{\Delta 181}$  mutants. (A,B) Pan-neuronal expression of ilp5 (C155-GAL4;  $mon1^{\Delta 181}$ >UAS-ilp5) rescues ovary size in  $mon1^{\Delta 181}$ mutants (compare A with B). (C,D) Expression of ilp5 in IPCs using ilp2-GAL4 (mon1 $^{\Delta 181}$ , ilp2-GAL4/mon1 $^{\Delta 181}$ , UAS-ilp5) rescues ovary size in mon1 mutants (compare C with D). (E) A comparison of the percentage distribution of egg chambers between control (C155-GAL4;  $mon1^{\Delta 181}$ ) and mutants expressing UAS-Mon1:HA, UAS-ilp5 and UASilp3. A significant increase in the number of egg chambers at stage 9-10 is observed upon expression of UAS-mon1:HA (mutant control: 1.37±0.59 versus 8.07±1.04 in C155-GAL4; mon1<sup>Δ181</sup>>UAS-mon1:HA). A smaller increase is seen with UAS-ilp5 (3.82±0.81) and UAS-ilp3 (5.15±1.61). A significant increase in the number of eggs is seen in all three genotypes ( $mon1^{\Delta 181}$ : 0 versus 9.07±1.28 in C155-GAL4; mon1<sup>Δ181</sup>>UAS-mon1:HA versus 4.9±2.19 in C155-GAL4;  $mon1^{\Delta 181}$ >UAS-ilp5 versus 6.11 $\pm$ 0.99 in C155-GAL4; mon1<sup>Δ181</sup>>UAS-ilp3). (F) Distribution of egg chamber stages in mutants expressing UAS-ilp5 and UAS-ilp3 in the IPCs. Note the increase in the number of stage 9-10 egg chambers [1.47±1.21 (mutant control) versus 5.95±0.93 (ilp2-GAL4; mon1<sup>Δ181</sup>>UAS-ilp5) versus  $2.3\pm0.94$  (ilp2-GAL4; mon1 $^{\triangle181}$ >UAS-ilp3)] and mature eggs [0 (mutant control) versus 7.78±0.98 (ilp2-GAL4; mon1<sup>Δ181</sup>>UAS-ilp5) versus 4.14±2.76 (ilp2-GAL4;  $mon1^{\Delta 181} > UAS-ilp3)]. *P<0.05; **P<0.01; ***P<0.001;$ Student's t-test was used to compare intensity of ILP5 staining. Egg chamber distribution (E,F) was analyzed using two-way ANOVA. N=number of animals, n=number of ovarioles.

Given that the sterility and ovarian phenotypes seen in mon1 mutants are reminiscent of those seen in *chico* mutants that have impaired insulin signaling, we checked for a possible connection between mon1 and insulin signaling. Two findings are of interest in this context. First, levels of a number of ILP genes (ilp1, ilp3, ilp5) and ilp6) are significantly reduced in  $mon1^{\Delta 181}$  mutants. Second, feeding an insulin-rich diet to homozygous  $mon1^{\Delta 181}$  females ameliorates the ovarian phenotypes induced by the loss of mon1 (Fig. 4).

Our data show that knockdown of *mon1* in the OPNs regulates ILP5 levels in the IPCs, pointing towards a non-cell-autonomous mode of regulation of ILP levels (Fig. 5). Furthermore, our results from the GRASP experiments show that OPNs make direct synaptic connections with the IPCs, suggesting that the regulation by Mon1 is likely to be direct between the two cell types (Fig. 5). Taken together, our results suggest that Mon1 in

OPNs is required to control production and/or secretion of ILPs in the IPCs.

The octopamine receptor OAMB is expressed in the IPCs and its knockdown has been shown to increase *ilp3* mRNA levels with no effect on *ilp2* and *ilp5* in the IPCs (Luo et al., 2014). This is in contrast to *mon1* mutants that show reduced levels of *ilp3* and *ilp5*, but not *ilp2*. The mechanism by which Mon1 regulates ILP expression and whether the circulating levels of the respective ILPs are also affected in a consistent manner still remains to be determined.

Our genetic analysis indicates that expression of ILP2 in the IPCs is not sufficient to rescue the lethality and ovarian phenotype in *mon1* mutants. Consistent with this, we found ILP2 levels to be elevated (Fig. S3), possibly in response to low levels of other ILPs. Altogether, our results argue that *mon1* functions in the OPNs in a manner that places it genetically upstream of insulin signaling to regulate ovary

maturation. Elucidating the mechanistic underpinnings of this regulatory relationship will be the focus for future studies.

It is remarkable that, in context of the neuromuscular junction, *mon1* appears to be secreted by the presynaptic terminals (Deivasigamani et al., 2015). This observation suggests that it is competent to function in a non-cell-autonomous manner. As in the case of the neuromuscular junction, it is possible that Mon1 engineers the expression of OAMB or other octopamine receptors, thereby indirectly influencing the activation of IPCs. Alternatively, Mon1 may facilitate the release of as yet unknown factors that influence expression of ILPs. These possibilities need to be explored by directly testing the levels and/or activity of OAMB receptors and ILPs in the IPCs.

Secretion of ILPs is thought to be regulated by ecdysone and juvenile hormone. The IPC/JH axis is part of an integrated network that regulates spatiotemporal growth and development in the fly. The interdependence of signaling of elements in the network, the presence of positive- and negative-feedback loops and the context dependence suggests that both the JH and DILPs are important players in the pre-vitellogenic checkpoint. We therefore tested whether Mon1 exerts its influence on oogenesis via these pathways. Arguing against the possibility, however, feeding  $mon1^{\Delta I81}$  mutant flies with 20-OH ecdysone did not rescue the ovarian phenotype to any appreciable extent. Topical addition of JH led to a mild rescue of the  $mon1^{\Delta 181}$  phenotype to about 15% of that of wild-type flies. In contrast, feeding excess levels of insulin effectively rescued the  $mon1^{\Delta 181}$ -dependent ovarian phenotype to about 70% of wild type. Furthermore, the extent of rescue could be correlated with the amount of insulin intake and the length of feeding time. For example, mutant flies fed on insulin-containing medium for 10 days show a near complete rescue of the ovarian phenotypes, including rescue of the ovariole number and presence of mature eggs. A similar rescue is seen at day 5, although these ovaries tend to have a few 'mutant-like' small ovarioles. Taken together, these observations strongly suggest that defects induced by the loss of Mon1 are, at least in part, due to impaired insulin signaling. Consistently, phospho-4EBP1 levels for whole-fly lysates are  $\sim$ 45% reduced in  $mon1^{\Delta 181}$  mutant flies (Fig. S4).

OPNs, to date, have not been implicated in ovarian development. Our data suggest that ILP3 and ILP5, but not ILP2 regulate the vitellogenic checkpoint and the production of ILPs in IPCs is, in turn, regulated by an unknown signal emanating from the OPNs. Mon1 in OPN neurons appears to function as a crucial node for activating this signal and thus regulates ILP3/ILP5 levels, adding an additional level of control for insulin signaling under normal feeding conditions, i.e. in the absence of starvation.

Our data reveal a novel regulatory loop involving a specific subset of neurons and insulin metabolism that, in turn, results in phenotypic abnormalities at the organismal level. Although the precise identity of the individual effectors is unclear at present, these data clearly demonstrate how systemic level communication between individual cells, tissues and even organs can participate in, and contribute to, organismal homeostasis.

#### **MATERIALS AND METHODS**

#### Fly husbandry and strains used

All stocks were reared on standard corn meal agar medium.  $mon1^{\Delta 181}$  and  $mon1^{\Delta 129}$  were generated through excision of pUAST-Rab21::YFP insertion using standard genetic methods (Deivasigamani et al., 2015). Df(2L)9062, C155-GAL4, tdc2-GAL4 (#9313), htl-GAL4 $^{\rm GMR93H07}$  (#40669) and nos-GAL4 are from the Bloomington Stock Center. The mon1 RNAi line (GD7852) is from the VDRC stock center. Other stocks were as follows:

*UAS-mon1::HA* (a kind gift from T. Klein, University of Dusseldorf, Germany), *ilp2*-GAL4 (Rulifson et al., 2002), *UAS-ilp5* and *UAS-ilp3* (G. Hasan, NCBS, Bangalore; Ikeya et al., 2002); *tdc2-LexA::p65* (#52242, Bloomington Stock Center); *UAS-nSyb-spGFP1-10*, *LexAop-CD4-spGFP11* (# 64314, Bloomington Stock Center); and *10XUAS-IVS-mCD8::RFP*, *13X LexAop2-mCD8::GFP* (# 32229, Bloomington Stock Center). Except when stated, all crosses were carried out at 25°C.

#### **Immunostaining and imaging**

For whole-ovary imaging, adult ovaries were dissected in 1× PBS at 4°C, and imaged using an Olympus Axio Zoom Stereoscope. All images were taken at the same magnification. For immunostaining, adult ovaries were dissected in 1× PBS at 4°C and fixed with 4% paraformaldehyde in 1× PBS with 0.3% Triton-X for 25 min at room temperature. The samples were washed six times in 1× PBS with 0.3% Triton-X for 15 min each and blocked in 2% BSA and 0.3% Triton-X in 1× PBS for 1 h. Incubation with primary antibody was carried out overnight at 4°C. Post-primary antibody incubation, the samples were washed four times for 10 min in 1× PBS containing 0.3% Triton-X and incubated with the appropriate fluorescent secondary antibody for 1 h at room temperature in the dark. The ovaries were then washed (six times for 5 min each in PBST) and mounted using 70% glycerol containing n-propyl gallate. A pair of ovaries from a single adult female was mounted per coverslip to count the number of ovarioles and to determine the proportion of different egg chambers per ovariole. The concentrations of the different dyes and antibodies were as follows: DAPI [Thermo-Fisher (Molecular Probes), D1306, 1:500]; phalloidin 488 [Thermo-Fisher (Molecular Probes), A12379, 1:100]; anti-Orb (DSHB, 1:20); Alexa Fluor secondary antibodies [Thermo-Fisher (Molecular Probes), A11031, A11034; 1:1000]. Imaging was carried out using the Leica SP8 confocal microscope system with a 40× oil objective (1.3 N.A.). Immunostaining of adult brains was carried out as described previously (Pfeiffer et al., 2010). Anti-ILP5 and anti-ILP2 (1:800 and 1:400 respectively; kind gifts from P. Leopold, Institut Curie, France). Image processing and intensity measurements were done using ImageJ software (NIH). The image of the ovary in Fig. 1H was created from two overlapping pictures and assembled by merging them in Photoshop to obtain the image of the entire ovariole. For the GRASP experiment (Fig. 5I), GFP staining was carried out using anti-chicken GFP (Thermo-Fisher, A10262, 1:1000). Identical staining was obtained with anti-rabbit GFP (Thermo-Fisher, A11122, 1:1000; Kim et al., 2012; Zhou et al., 2015) and anti-RFP (a kind gift from R. Hegde, Cambridge, UK; 1:2000). GraphPad Prism was used for statistical analyses. All data are presented as mean±s.e.m. Figures were assembled using Adobe Photoshop CS4.

#### **Quantitative RT-PCR**

Estimation of orb levels was carried out using quantitative PCR on cDNA synthesized from mRNA isolated from adult wild-type (w1118) and mon1 mutant ovaries. The ovaries were dissected in ice-cold 1× PBS. RNA isolation was carried out using Trizol reagent according to the manufacturer's instructions. For estimation of ILPs, mRNA was isolated from 1- to 2-day-old adult female flies. In each case, 1 µg of RNA was used for the reverse transcription reaction. The sequence of the primers used for quantitative PCR are as follows: rp49-f, GACGCTTCAAGGGACAGTATC; rp49-r, AAAC-GCGGTTCTGCATGAG; orb-f, GAGTGGGAAAGGGAAGGTAC; orb-r, CAGGTAGGAGCTTATCGAGTG; ilp1-f, GGTGTGTCCCCATGGCT-TTA; ilp1-r, TGCTGCTATCATCCTGCACC; ilp2-f, CGAGGTGCTG-AGTATGGTGTG; ilp2-r, CCCCAAGATAGCTCCCAGGA; ilp3-f, ATG-GGCATCGAGATGAGGTG; ilp3-r, CGTTGAAGCCATACACACAGAG; ilp4-f, ATGAGCCTGATTAGACTGGGAC; ilp4-r, TCTAGCATCCTTA-GACGCACT; ilp5-f, TGCCTGTCCCAATGGATTCAA; ilp5-r, GCCAAG-TGGTCCTCATAATCG; ilp6-f, GTCCAAAGTCCTGCTAGTCCT; ilp6-r, TCTGTTCGTATTCCGTGGGTG; ilp7-f, AGGAGGGTCTCGAGATG-CTT; ilp7-r, CCCAATATAGCTGGCGGACC; ilp8-f, GGACGGACG-GGTTAACCATT; and ilp8-r, CATCAGGCAACAGACTCCGA.

#### Feeding experiments

Regular corn-meal medium was supplemented with  $5\,\mu g/ml$  of bovine insulin (Sigma-Aldrich, I6634). Newly eclosed wild-type and  $monl^{\Delta I81}$ 

mutant females were transferred to insulin-containing medium. Ovaries were examined in 5-day-old animals. For measurement of life span, the animals were maintained on insulin medium for the entire duration of the experiment. For comparison of feeding behavior of  $mon1^{\Delta 181}$  with wild-type flies, we used a food dye (Brilliant Blue FCF, Sigma 80717). Animals are starved for 4 h and fed on either 0.5% sucrose-soaked filter paper or cornmeal agar for 1 h. In both cases, 0.5% dye is incorporated in the nutrient medium. The amount of dye intake was measured by spectrophotometry ( $\lambda$ 629, Varioscan plate reader) for eight animals (N=3 experiments) in fly lysates that were homogenized in water and centrifuged for 15 min at 14,000 g. Two-day-old  $mon1^{181}$  adults appeared to ingest equal amounts of food/sucrose when compared with age-matched  $mon1^{181}$ /+ or  $w^{1118}$  adults.

#### **Application of Juvenile hormone**

Acetone was used as a vehicle to dissolve and apply Juvenile hormone (JH) III (J2000, Sigma). One-day-old adult flies were topically administered JH by pipetting 0.5  $\mu$ l of solution onto the abdomen of the fly. JH was applied at the concentrations of 0.1, 1 and 5  $\mu$ g per animal. The concentrations used were consistent with the range used in earlier studies (Gruntenko et al., 2019; Rauschenbach et al., 2014; Terashima and Bownes, 2004). The animals were kept on yeasted corn-meal agar for 48 h before dissection of ovaries and counting of the number of eggs (stage 13-14) per animal. There was an increase in the number of eggs (stage 13-14) from <1 per animal ( $mon1^{\Delta I8I}$ ) to  $4.77\pm2.0$  eggs per animal in response to JH application when compared with wild-type animals (33±3.2 eggs per animal).

#### Analysis of egg chamber distribution

All analysis to determine the distribution of egg chambers was carried out on 2- to 3-day-old virgin females maintained on regular cornmeal agar medium containing yeast soon after eclosion. The number of egg chambers for each stage were scored on a 'per animal' basis. Except where stated, the data presented for each stage is an average for 5 animals (n=5).

#### **Fecundity assay**

Individual newly eclosed virgin females of the appropriate genotype were crossed to single  $w^{III8}$  males and transferred on the same day to a sucrose agar medium containing a small amount of yeast paste. Sucrose plates were changed every 24 h. The day of eclosion was treated as 'day 0'; egg laying was monitored from the 2nd day post-eclosion.

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#### Competing interests

The authors declare no competing or financial interests.

#### Author contributions

Conceptualization and methodology: A.R.; Investigation: N.D., A.R., K.S.,S.T.; Formal analysis: N.D., A.R., K.S., S.T., G.D., G.S.R., Visualization: A.R.; Resources: A.R., G.S.R.; Supervision: A.R.; Data curation: N.D., A.R.; Writing original draft: A.R., G.D.; Writing review and editing: A.R., G.S.R and G.D; Project administration: A.R. and G.S.R; Funding acquisition: A.R., G.S.R.

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#### Supplementary information

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