Neuronal Control of Metabolism through Nutrient-Dependent Modulation of Tracheal Branching

Citation for published version:
Linneweber, GA, Jacobson, J, Busch, KE, Hudry, B, Christov, CP, Dormann, D, Yuan, M, Otani, T, Knust, E, de Bono, M & Miguel-Aliaga, I 2014, 'Neuronal Control of Metabolism through Nutrient-Dependent Modulation of Tracheal Branching', Cell, vol. 156, no. 1-2, pp. 69-83.
https://doi.org/10.1016/j.cell.2013.12.008

Digital Object Identifier (DOI):
10.1016/j.cell.2013.12.008

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Cell

Publisher Rights Statement:
This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

General rights
Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.
Neuronal Control of Metabolism through Nutrient-Dependent Modulation of Tracheal Branching

Gerit A. Linneweber,1 Jake Jacobson,1 Karl Emanuel Busch,2,6 Bruno Hudry,1 Christo P. Christov,3 Dirk Dormann,1 Michaela Yuan,1 Tomoki Otani,3,6 Elisabeth Knust,4 Mario de Bono,2 and Irene Miguel-Aliaga1,*

1Gut Signalling and Metabolism Group, MRC Clinical Sciences Centre, Imperial College London, Du Cane Road, London W12 0NN, UK
2MRC Laboratory of Molecular Biology, Francis Crick Avenue, Cambridge CB2 0QH, UK
3Department of Zoology, University of Cambridge, Downing Street, Cambridge CB2 3EJ, UK
4Max Planck Institute of Molecular Cell Biology and Genetics, Pfotenhauerstrasse 108, 01307 Dresden, Germany
5Present address: Centre for Integrative Physiology, The University of Edinburgh, Hugh Robson Building, Edinburgh EH8 9XD, UK
6Present address: The Gurdon Institute, University of Cambridge, Tennis Court Road, Cambridge CB2 1QN, UK
*Correspondence: i.miguel-aliaga@imperial.ac.uk
http://dx.doi.org/10.1016/j.cell.2013.12.008
This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

SUMMARY

During adaptive angiogenesis, a key process in the etiology and treatment of cancer and obesity, the vasculature changes to meet the metabolic needs of its target tissues. Although the cues governing vascular remodeling are not fully understood, target-derived signals are generally believed to underlie this process. Here, we identify an alternative mechanism by characterizing the previously unrecognized nutrient-dependent plasticity of the Drosophila tracheal system: a network of oxygen-delivering tubules developmentally akin to mammalian blood vessels. We find that this plasticity, particularly prominent in the intestine, drives—rather than responds to—metabolic change. Mechanistically, it is regulated by distinct populations of nutrient- and oxygen-responsive neurons that, through delivery of both local and systemic insulin- and VIP-like neuropeptides, sculpt the growth of specific tracheal subsets. Thus, we describe a novel mechanism by which nutritional cues modulate neuronal activity to give rise to organ-specific, long-lasting changes in vascular architecture.

INTRODUCTION

Unlike the more stereotypical development of the body’s main blood vessels, the formation of the capillary networks responsible for tissue perfusion is an adaptive process primarily governed by the metabolic needs of the target tissues (Fraisil et al., 2009; Potente et al., 2011). The plastic nature of this adaptive angiogenesis is further highlighted by the dramatic changes in vascularization observed in tumors or in obese adipose tissue: changes that contribute to the progression of pathologies such as cancer and obesity and are becoming increasingly central to their treatment (Cao, 2010; Kerbel, 2008; Lijnen, 2008). Although environmental factors such as diet are widely believed to affect the development and progression of these pathologies, exploration of the link between nutrition and angiogenesis has largely been confined to correlative studies. These include descriptions of the effects of gestational nutrition on the placental vasculature (Belkacemi et al., 2010; Rutland et al., 2007) or the pro/anti-angiogenic actions of nutrients and metabolites with a potential modulatory role in cancer (Adolphe et al., 2010; Kumar et al., 2013). A tantalizing new study has shown that increasing adipose tissue vascularization can ameliorate the deleterious metabolic effects of a high-fat diet, pointing to a central metabolic role for these vascular changes (Sung et al., 2013). However, whether modulation of angiogenesis is associated with metabolic benefits remains a controversial topic, partly because it is not trivial to genetically target the blood vessels of specific organs to recapitulate the changes associated with certain dietary interventions without affecting other cell types or vascular pools (Cao, 2010; Lijnen, 2008; Sun et al., 2012; Sung et al., 2013). Regardless of its metabolic consequences, adaptive angiogenesis is widely believed to be mechanistically driven by target-derived signals (Cao, 2007; Fraisil et al., 2009).

A close spatial association between mammalian nerves and vessels was observed as long ago as 1543 (Vesalius, 1543), an association that has subsequently been shown to result from mutual guidance or common pathfinding mechanisms during the formation of the neural and vascular networks (Carmeliet and Tessier-Lavigne, 2005; Mukoyama et al., 2005; Mukoyama et al., 2002; Quaegbeur et al., 2011). Notably, interplay of innervation and vascularisation of internal organs has also been described (Davies, 2009). A functional role for these neurovascular interactions was suggested following the discovery that vessel abnormalities precede a number of neurodevelopmental and neurodegenerative disorders: an observation that points to
angiogenesis as a therapeutically relevant process (Quaegebeur et al., 2011; Storkebaum et al., 2011). The question remains whether, in a reciprocal process, neuronal activity may affect adaptive angiogenesis. In spite of some intriguing associations (Asano et al., 1997; Tonello et al., 1999), no neuronal populations have been identified that effect long-lasting changes in angiogenesis in response to environmental factors.

Drosophila melanogaster has an open circulation, but its tracheal system has a role analogous to that of the vertebrate vasculature in supplying tissues and internal organs with oxygen (Fraisil et al., 2009; Uv et al., 2003). During embryogenesis, developmental mechanisms akin to those discovered in the vertebrate lung and vasculature make use of signaling pathways such as fibroblast growth factor (FGF) signaling to sculpt this complex tracheal network of interconnected tubes (Ghabrial et al., 2003; Javerzat et al., 2002; Metzger et al., 2008; Uv et al., 2003). These embryonic proliferative and morphogenetic stages are superseded by a larval period of extensive, but mechanistically less understood, cellular growth. Growth is particularly prominent in the tracheal terminal cells: the cells at the end of each airway that make contact with target tissues and through which gas exchange takes place (Ghabrial et al., 2003; Uv et al., 2003). Like vertebrate capillaries, Drosophila tracheal terminal cells branch profusely in response to low oxygen using conserved FGF and hypoxia-inducible factor (HIF) signaling pathways (Centanin et al., 2008; Jarecki et al., 1999). This hypoxic remodeling has been assumed to be the only source of tracheal plasticity and, in normal conditions, the tracheal system is generally believed to grow in proportion to the whole organism. In this study, we use a combination of genetic approaches, metabolic profiling, and in vivo imaging to uncover previously unrecognized nutritional plasticity in the fly tracheal system. In contrast to the known target-derived mechanisms of adaptive remodeling, we find this plasticity to be regulated by a mechanism, previously undescribed in either flies or vertebrates, involving nutrient-responsive neurons effecting long-lasting and metabolically significant changes in tracheal architecture.

RESULTS

Branching of Tracheal Terminal Cells Is Regulated in an Organ-Specific Fashion According to Both Previous and Current Nutrient Availability

While subjecting wild-type Drosophila larvae to different dietary conditions, we observed that a severe reduction in dietary yeast (the main source of lipid and amino acids in the larval diet) was accompanied by an almost ubiquitous reduction in tracheal terminal cell branching, even when controlling for overall developmental delay by allowing nutrient-restricted larvae to develop to a comparable stage (Figures 1A–1C, 1G–1I and Figures S1A, S1C, S1D, S1F, S1G, S1I, S1J, S1L, S1M, and S1O available online). The single exception was the tracheal branches of the central nervous system (CNS), which were refractory to this dietary manipulation (Figures 1A and 1G). By contrast, a mild reduction in dietary yeast neither affected developmental timing nor led to major changes in the size of organs or that of most tracheal terminal cells (Figures 1D and 1E and data not shown) but did lead to a severe reduction in tracheal coverage throughout the digestive tract (Figures 1F, S1B, S1E, S1H, S1K, and S1N). Reduced tracheal coverage was not caused by cell death (Figures S2A and S2B) and could not be solely accounted for by defects in gas filling (Figures S2Q and S2R). Instead, it resulted from reduced tracheal terminal cell branching (Figures S2E–S2Q, S2K–S2M, and S2Q–S2R). To investigate the reversibility of the tracheal changes described above, we reared larvae under the mild nutrient restriction conditions shown to reduce intestinal tracheation and transferred them to more nutritious food immediately after eclosion. Even after 7 days on a nutritious diet, the intestinal tracheae of these adults flies were significantly less branched than those of control adult flies always reared on a nutritious diet (Figures 1J and 1K), indicating that a defined period of nutrient restriction has long-term effects on the tracheal scaffold.

Dietary plasticity could be a feature unique to larval tracheae, given that their branches are undergoing extensive growth. To investigate whether adult tracheae are also responsive to diet, we allowed wild-type flies to develop under our standard nutritional conditions and then exposed them to nutritionally poor or imbalanced diets as adult flies. As Figure 1L shows, a 7 day nutritional imbalance (9% sucrose) led to increased intestinal tracheation of the mid-midgut, confirming the dietary plasticity of the tracheal system also in adult flies.

Collectively, these data uncover previously unrecognized nutritional plasticity of the insect tracheal system, shaped by both previous and current nutritional states. The tracheae of different organs exhibit different degrees of nutritional plasticity; intestinal tracheal terminal cells are particularly sensitive to a reduction in yeast availability, while CNS tracheae are preferentially spared.

Differential, Cell-Autonomous Activation of Insulin Signaling Mediates Tracheal Terminal Cell Growth and Underlies the Enhanced Plasticity of Intestinal Tracheae

We then focused on the larval phenotypes to investigate the molecular mechanisms of nutritional plasticity. Hypoxia, the only known regulator of tracheal plasticity, has been shown to promote tracheal branching by inducing FGF ligand in target tissues and receptor upregulation in tracheal cells (Centanin et al., 2008; Jarecki et al., 1999). Although downregulation of the FGF receptor gene Breathless (Btl) did lead to reduced tracheation in most scored tissues, consistent with the known FGF requirement for the establishment of the tracheal scaffold during earlier developmental stages (Ghabrial et al., 2003; Uv et al., 2003), further attempts to manipulate FGF signaling or to detect FGF ligand expression and differential pathway activation under different nutritional conditions all failed to support a role for FGF signaling in coupling nutrition with larval tracheal growth (data not shown). These included expression of Btl, constitutively active Btl, and its ligand Branchless (Bnl) in tracheal terminal cells and analysis of Bnl and Stumps (a downstream signaling component) expression under different nutritional conditions. We then turned our attention to the insulin signaling pathway: the major coordinator of nutrient intake and tissue size in all animals including Drosophila (Andersen et al., 2013). We first suppressed the intracellular insulin signal transducer phosphoinositide 3-kinase (PI3K) by expressing the dominant-negative
Dp110D954A (referred to as PI3K-DN; Leevers et al., 1996) in tracheal terminal cells using DSRF-GAL4 (Gervais and Casanova, 2011). This led to reduced tracheal terminal cell branching both in the periphery and throughout the digestive tract, but not in the CNS (Figures 2A–2F, S2S, S2T, S3A–S3D and data not shown): a reduction qualitatively and quantitatively comparable...
Figure 2. Organ-Specific Effects of Reduced Insulin Signaling on Tracheal Coverage

(A–C) Representative tracheation of the areas boxed in the cartoons in control larvae: body wall (A), midgut (anterior, B), and hindgut (mid-hindgut, C).

(D–F) Expression of PI3K-DN in tracheal terminal cells leads to reduced branching in body wall (D), midgut (anterior, E), and hindgut (mid-hindgut, F). For body wall: p = 0.001 (DSRF>PI3K-DN versus GAL4 control), p < 0.001 (DSRF>PI3K-DN versus UAS control), n = 13–15/set. For anterior midgut: p < 0.0001 (DSRF>PI3K-DN versus GAL4 control), p < 0.001 (DSRF>PI3K-DN versus UAS control), p = 0.03 (GAL4 versus UAS control), n = 15/set. For mid-hindgut: p < 0.0001 (DSRF>PI3K-DN versus GAL4 control), p < 0.0001 (DSRF>PI3K-DN versus UAS control), n = 19–25/set.

(legend continued on next page)
to that observed in these tracheal terminal cells following severe nutrient restriction (Figures 1G–1I). As in the case of diet, the intestinal branches appeared to be more severely affected by this manipulation. Because the selective intestinal phenotype was not caused by stronger GAL4 expression in intestinal tracheae (data not shown), we tested whether it resulted from increased sensitivity to insulin signaling. To this end, we made use of an RNAi construct against the insulin receptor (InR) known to lead to incomplete receptor downregulation and a milder reduction in insulin signaling (Sialdina et al., 2009; Willecke et al., 2011). Driving this RNAi transgene in all tracheal terminal cells led to a significant reduction in intestinal, but not body wall or CNS, tracheal coverage (Figures 2G–2I, S3E, and S3F and data not shown). As in the case of dietary or PI3K manipulations, reduced coverage resulted from reduced tracheal terminal cell branching (Figures S2C, S2D, S2H–S2J, and S2N–S2P).

Together, these results confirm the cell-autonomous role for the insulin signaling pathway in the regulation of tracheal terminal cell growth and suggest that the enhanced nutritional plasticity of the gut tracheae is a consequence of their higher sensitivity to insulin signaling.

Different Tracheal Subsets are Combinatorially Modulated by Both Systemic and Local Insulin- and VIP-like Neuropeptides

In Drosophila larvae, nutrient restriction leads to growth inhibition, caused by the reduced release of several insulin-like peptides (Ilps) from brain insulin-producing cells (the so-called median neurosecretory cells, mNSCs, represented schematically in Figure 3J) into the hemolymph (Géminard et al., 2009). A triple mutation of the three main mNSC Ilps (Ilp2, Ilp3, and Ilp5; Grönke et al., 2010) largely recapitulated the phenotype resulting from expression of PI3K-DN in tracheal terminal cells. Indeed, reduced growth was observed in both body wall (Figures 3A and 3D) and intestinal tracheal terminal cells (Figures 3B, 3E, and S4A–S4D). However, we found the posterior hindgut tracheal branches to be spared in these larvae (Figures 3C and 3F). Immunohistochemical and ultrastructural analyses of this intestinal portion revealed that these posterior tracheal branches were adjacent to the two hindgut nerves that run along both sides of the hindgut (Figures 3K, 3L, and 3B). We have previously shown that axons emanating from a different population of CNS Ilp-producing neurons, the Ilp7 neurons, contribute to this innervation (Figure 3J; Miguel-Alia et al., 2008) and thus could provide a local peptide supply to this portion of the gut. Functional inactivation of the Ilp7 neurons by expression of the inward-rectifying potassium channel Kir2.1 or by expression of tetanus toxin light chain did not affect most tracheae but led to reduced tracheal coverage of two portions of the hindgut (Figures 3G–3I, S4E, and S4F and data not shown): the posterior hindgut (Figures 3C and 3I), where the Ilp7 axons are adjacent to the posterior visceral tracheal branches (Figures 3K and 7B), but also the mid-hindgut (Figure S4F), which we had also found to be regulated by systemic mNSC-derived Ilps (Figures S4B and S4D). In this latter region, the visceral tracheal branches emanate from the segmentally repeated main lateral branches (Figures 7A and 7B) and do not abut the Ilp7 axons, suggesting paracrine growth regulation.

We then characterized the peptidergic profile of the central neurons contributing to the hindgut nerves using immunohistochemistry. Four of the eight Ilp7-expressing neurons coexpress pigment dispersing factor (Pdf) (Figures 4A and 4B): a neuropeptide that shares functional and signaling similarities with vertebrate vasoactive intestinal polypeptide (VIP) (Taghert and Nitabach, 2012). Four other central hindgut-innervating neurons also express Pdf and bundle together with the Ilp7 hindgut nerves (Figure 4B; Talsma et al., 2012). Together, both neuronal populations deliver Pdf and Ilp7 to the hindgut in a regionalized fashion: Ilp7 is apparent only in the posterior hindgut, whereas Pdf is present in both posterior and mid-hindgut terminals (Figures 4C and 7B). Mutation of these peptides, alone or in combination, revealed complex control of different intestinal tracheal subsets by local Ilp7 and Pdf peptides in combination with the systemic Ilp2, Ilp3, and Ilp5 peptides (Figures 4D–4U, S5A–S5L, and 7B): in the posterior hindgut, neither loss of Ilp7 alone nor Ilp2, Ilp3, and Ilp5 together affected tracheal branching (Figures 4F, 4I, S5C, 3C, and 3F), but loss of all four peptides resulted in reduced tracheal terminal cell growth (Figures 4L and S5C), indicating partially redundant control of tracheal terminal growth. Loss of Ilp7 or Pdf alone, or tracheal-specific downregulation of the Pdf receptor (DSRF-GAL4, UAS-Pdfr-RNAi), resulted in reduced tracheal growth only in the mid-hindgut (Figures 4G–4I, 4M–4R and S5A–S5I); a region also affected by the lack of systemic Ilps (Figures S4B and S4D) and not directly exposed to Ilp7 peptide (Figure 7B). Finally, mutants lacking both Ilp7 and Pdf displayed reduced tracheal growth in both the mid-hindgut and posterior hindgut (Figures 4T, 4U, S5K, and SSL), indicating that Ilp7 and Pdf act redundantly in the posterior hindgut.

Collectively, neuropeptide mutation and tracheal receptor downregulation experiments indicate that growth of tracheal terminal cells is directly regulated by the nervous system. The systemically secreted Ilps act as virtually pan-tracheal regulators, but in some intestinal portions they synergize in a combinatorial—and sometimes partially redundant—manner with locally delivered Ilp and Pdf neuropeptides.

Exposure to Nutrients and Reductions in Oxygen Availability Elicit Calcium Responses in the Gut-Innervating Ilp7/Pdf Neurons

Both mNSCs and Ilp7 neurons have been shown to modulate feeding responses to nutrient scarcity in adult flies (Cognigni et al., 2011). However, the dietary dependency of Ilp release has only been investigated in mNSCs using immunohistochemistry (Géminard et al., 2009). To directly image neural activity in...
Figure 3. Two Subsets of Insulin-Producing Neurons Regulate the Growth of Different Tracheal Subsets

(A–C) Representative terminal tracheation in well-fed control larvae. The specific body wall/gut areas are boxed in the cartoons: body wall (A), midgut (B, anterior), and hindgut (C, posterior).

(legend continued on next page)
response to nutrients in vivo, we expressed the genetically encoded green fluorescent Ca\(^{2+}\) indicator GCaMP3 (Tian et al., 2009) in Ilp7 neurons, together with a red fluorescent protein to visualize the cell bodies. Ilp7 cell bodies displayed some transient activity in the absence of a stimulus, which rapidly increased following yeast presentation (Figures 5A, 5C, S6A, and Movie S1). In most neurons, the frequency and amplitude of the transient Ca\(^{2+}\) peaks increased and then adapted after about one minute, possibly a consequence of persistent exposure to yeast. This response was yeast-specific because exposure to sucrose did not elicit any responses in these neurons (data not shown), consistent with the yeast dependency of tracheal growth. It was also specific to Ilp7 neurons, given that GCaMP3 fluorescence intensity was unaffected by yeast in the Capa-expressing Va neurons, used as a control population of six unrelated peptidergic effenter neurons (Suska et al., 2011) (Figure 5A).

The only well-characterized environmental trigger of tracheal branching is hypoxia (Centanin et al., 2008; Jarecki et al., 1999). We therefore monitored oxygen-evoked Ca\(^{2+}\) responses in these two neuronal populations and found that hypoxia led to a fast and very robust response in the Ilp7—but not in the Va—neurons (Figures 5B, 5D, and S6B, and Movie S2). This response was qualitatively distinct from that resulting from yeast exposure. Indeed, it was predominantly tonic, although some animals mainly showed transient peaks of increased amplitude, and lasted throughout the hypoxic period, decreasing slightly over time. Interestingly, the return to normoxia almost completely abrogated the basal transient firing of Ilp7 neurons, suggesting hyperpolarization. This effect was not a consequence of excessive firing and cellular “exhaustion” because repeated hypoxic stimulation continued to activate the Ilp7 neurons (Figure S6C).

Together, these findings indicate that the activity of the Ilp7- and Pdf-producing neurons is increased in vivo by both nutritional cues and reductions in oxygen availability.

**Activation of the Ilp7/Pdf Neurons Promotes Tracheal Branching Locally**

Together with previous Ilp/Pdf loss-of-function experiments, the above experiments suggested that nutritional modulation of Ilp neuronal activity underlies the nutritional plasticity of tracheae. To test this idea, we used thermogenetics to achieve persistent, low-level activation of the Ilp7 neurons throughout larval life by expressing the heat-sensitive channel TrpA1 from Ilp7-GAL4 in larvae reared at 25°C. This promoted tracheal branching in a paracrine fashion; it increased branching of the adjacent visceral tracheal branch of the posterior hindgut and the tracheal terminal cells of the neighboring mid-hindgut, but did not redirect those of the anterior hindgut or other regions (Figures 5E and 5F, and data not shown). Hence, in addition to being necessary, Ilp7 neurons are sufficient to sustain tracheal growth in the hindgut.

**The Organ-Specific Modulation of Tracheation Is Metabolically, but Not Developmentally, Significant**

The finding that tracheal branching is directly regulated by nutrient-responsive neurons suggests that tracheal terminal cells may be used by the nervous system as effectors of metabolic adaptations to nutrient availability. To investigate this possibility, we recapitulated the differential effects of nutrient restriction on tracheae by either reducing tracheal terminal cell growth in all tissues (except for the CNS tracheae, using DSRF>btl-RNAi), or specifically in the gut tracheae (using DSRF > InR-RNAi). Reduced tracheation of all tissues did not affect larval development (Figure 6A) or carbohydrate metabolism (Figures S7A–S7C) but resulted in leaner larvae (Figure 6C) with reduced lipid stores (Figure 6D) and increased hemolymph glycerol (a metabolite derived from the hydrolysis of triglycerides) (Figure 6E), consistent with reduced lipid storage capacity in the fat body. These larvae did manage to eclose as adults but were sick and short-lived even in the presence of nutritious food (Figure 6F). By contrast, when reduced tracheation was confined to the gut, no developmental or metabolic phenotypes were apparent in larvae (Figures S7D–S7H), and there was no difference in adult lifespan between the experimental flies and controls on nutritious food (Figure 6F). We then hypothesized that the specific effect of nutrient restriction on gut tracheae may fulfill an adaptive role to allow flies to deal with poor nutritional conditions. To test this idea, we exposed the DSRF> InR-RNAi flies with reduced gut tracheation to a low-calorie diet throughout their adult lifetime, and found them to be significantly more resistant to nutrient scarcity than control flies: a tracheal phenotype that was confirmed using the recently published tracheal driver 14D03-GAL4 (Guo et al., 2013) (Figure 6G and data not shown). Metabolic profiling of these adult flies revealed no differences in carbohydrate metabolism but showed a reduction in lipid stores in poor nutritional conditions (Figures 6H, 6I, and S7I–S7L).

In summary, manipulations that recapitulate the effects of nutrient restriction and reduced insulin signaling specifically in...
tracheal terminal cells show that these cells are important metabolic mediators.

**DISCUSSION**

**Nutrient-Responsive Neurons as Effectors of Adaptive Tracheal Changes**

Our work has uncovered a new mechanism coupling nutrition and metabolism. In response to specific nutritional cues, small subsets of neurons are activated to regulate tracheal branching in an organ-specific and metabolically significant fashion. At least one of the two yeast-responsive neuronal subsets also responds to reduced oxygen—the other environmental modulator of tracheal branching in flies—so it will be interesting to determine the contribution of these neurons to the previously reported tracheal adaptations to hypoxia. Importantly, our identification of a shared neuronal substrate for both nutritional and hypoxic stimuli is, to our knowledge, the first of its kind in invertebrates and one remarkably similar to the mammalian carotid body: a cluster of chemoreceptors that monitors arterial oxygen concentration and nutrient levels to regulate breathing and cardiovascular tone (Pardal and López-Barneo, 2002; Prabhakar, 2000). Future work will aim to establish whether these *Drosophila* neurons are able to sense oxygen and/or nutrients directly and whether they do so using mechanisms akin to those described in the carotid body. This would lend further support to the existence of an evolutionarily conserved link between oxygen and nutrient neuronal sensing.

Molecularly, the neuronal control of different tracheal subsets involves both local and systemic actions of insulin- and VIP-like neuropeptides: neuronal mechanisms that are particularly complex and combinatorial along the digestive tract (Figure 7) and that differ from the known adaptive target-derived signals that sculpt tissue-specific angiogenesis (Cao, 2007; Fraisl et al., 2009). In this regard, tracheal cells can be seen as “metabolic motor neurons”; as the nervous system modulates motor neuron activity to regulate muscle contraction, it also modulates the branching of tracheal terminal cells to control the metabolic state of cells such as those of the fat body or the gut epithelium. It will be of interest to investigate whether similar mechanisms are deployed in vertebrates to effect long-lasting, tissue-specific and metabolically significant changes in angiogenesis in response to nutrition, in a manner distinct from (but reminiscent of) the acute changes in blood supply effected by neurons acting on blood vessel musculature (see, for example, Matheson et al., 2000).

**Organ-Specific Regulation of the Tracheal System by Local and Systemic Insulin- and VIP-like Neuropeptides**

In *Drosophila*, previous gain- and loss-of-function experiments had failed to reveal unique functions for most of the eight known Ilps (Brogiolo et al., 2001; Grönke et al., 2010). The regional regulation of tracheal subsets hence provides one possible explanation for the apparent redundancy of the *Ilp* gene family in *Drosophila*: while all these Ilps may indeed have the same function (in this case, to modulate tracheal growth in response to nutrition), they may carry it out in different places—for example, in the posterior hindgut in the case of Ilp7 and in other parts of the digestive tract for Ilp2, Ilp3, and Ilp5. This regional control of tracheal growth may extend to other regions: gut visceral musculature and CNS glia are known to activate Ilp3 and *Ilp2/Ilp6* gene expression respectively in a nutrient-dependent fashion (Chell and Brand, 2010; O’Brien et al., 2011; Sousa-Nunes et al., 2011). In light of our findings and the recent discovery that intestinal tracheae can regulate stem cell proliferation (Li et al., 2013), it is possible that local regulation of tracheal branching by Ilps contributes to their reported action on intestinal or neuronal stem cell proliferation (Chell and Brand, 2010; O’Brien et al., 2011; Sousa-Nunes et al., 2011).

Effects of insulin and VIP on blood vessels have been described in vertebrates (Chaudhuri et al., 2012; Holzer, 2006). Indeed, although the effect of Pdf on intestinal tracheal branching is unexpected in *Drosophila* (where this peptide is known for its central role in clock neurons; Taghert and Nitabach, 2012), neurally derived VIP has a vasodilatory effect on the arterioles of small intestine and colon (Holzer, 2006). However, the physiological significance of these (largely ex vivo) observations has not been entirely elucidated (Matheson et al., 2000). In contrast to this mode of regulation, involving acute modulation of endothelial muscle tone, the evidence for longer-lasting effects of these peptides on angiogenesis—which would be more akin...
to their action on the *Drosophila* tracheal system—is more tenuous and often contradictory (see, for example, Ogasawara et al., 1999; Ribatti et al., 2007). Our findings suggest that their effects may have been underestimated because they act in partially redundant fashion and in response to specific nutritional cues. Mechanistically, it has been proposed that the vertebrate peptides regulate proangiogenic target-derived signals. By contrast, our tracheae-specific receptor downregulation experiments clearly indicate that these peptides can act directly on the tracheal cells, so it will be of interest to establish whether both modes of action contribute to their effects on vertebrate angiogenesis.

**Metabolic Significance of the Tracheal Nutritional Plasticity**

In *Drosophila*, whole-organism manipulations of insulin signaling such as ablation of insulin-producing cells or Ilp mutation result in both slower development and "diabetic" phenotypes, highlighting their dual insulin/IGF-like role (Grönke et al., 2010; Rulifson et al., 2002). Strikingly, downregulation of insulin signaling only in one cellular target—the tracheal terminal cells—uncouples the developmental from the metabolic phenotypes of these peptides, thus identifying the tracheal system as an important and previously unrecognized metabolic target of insulin signaling in the fly. Hence, the tracheal involvement in previously reported

---

**Figure 5. Regulation of Ilp7 Neuronal Activity by Nutrients and Hypoxia, and Its Effect on Tracheal Branching**

(A) Exposure to yeast leads to a transient Ca$^{2+}$ rise in Ilp7 neurons. Activity returns to basal levels after one minute. No such response is observed in control Va neurons.

(B) A switch from 21% to 1% ambient O$_2$ elicits a rapid rise in Ca$^{2+}$ in Ilp7 neurons that persists while O$_2$ is low. Upon return to normoxia, the basal activity of the Ilp7 neurons is immediately abrogated. No Ca$^{2+}$ rise is triggered in control Va neurons, which display a subtle drop in Ca$^{2+}$ levels in response to hypoxia, as has previously been observed for different types of neurons in various species (Cheung et al., 2006; Fujimura et al., 1987; Krnjević, 1999). Error bars denote SEM.

(C and D) False color-coded single frames depicting GCaMP fluorescence in representative movies illustrating the response to yeast (C) or hypoxia (D) observed in Ilp7 neuronal cell bodies. Yellow/white indicates strong responses, red, low Ca$^{2+}$ (false color scale is shown to the left).

(E and F) 25°C thermogenetic activation of the TrpA1 channel expressed in Ilp7 neurons through larval development results in increased tracheal coverage of the midgut (F) relative to controls (E for GAL4 control). Quantifications are displayed to the right of these two panels (p < 0.001 versus GAL4 control, p < 0.0001 versus UAS control, n = 23-27/set).

Scale bars, 25 µm (C) and (D) or 10 µm (E) and (F). See also Figure S6.
A. Developmental time (larva)

B. Survival, well fed

C. Length/width (L3)

D. TAG/protein (L3)

E. Hemolymph free glycerol (L3)

F. Survival, well fed

G. Survival, nutrient restriction

H. TAG/protein (adult), well fed

I. TAG/protein (adult), nutrient restriction

(legend on next page)
insulin-modulated phenotypes, such as lifespan or resistance to oxidative stress (Grönke et al., 2010), deserves further investigation. Interestingly, a pan-tracheal reduction in insulin signaling results in normal carbohydrate metabolism but leads to reduced adiposity. This is suggestive of abnormal lipid metabolism in the fat body and is consistent with the recent finding that reduced fat mass in young mice (Sung et al., 2013) — although in both mice and flies this phenotype may eventually prove to be deleterious (Sung et al., 2013 and Figure 6E). Reduced adiposity is a phenotype that, although also consistent with one of the classic symptoms of type I diabetes in humans, had not previously been observed in flies with a ubiquitous reduction in insulin signaling or lacking the systemic Ilp peptides (puzzlingly, these flies were actually found to accumulate triglyceride; Böhni et al., 1999; Grönke et al., 2010). We suggest that this increased adiposity may have been secondary to the IGF-like effects of Ilps on developmental time, and only by uncoupling these developmental from the metabolic effects of Ilps, as we have done with the tracheal-specific reduction of insulin signaling, can some of the “true insulin-like” phenotypes of Ilps be unmasked.

We have also found that subtle changes in insulin signaling or in the nutritional content of the fly’s diet (some of which are within the range of those normally found in diets used for fly rearing in different labs) have a striking effect on an unexpected tracheal population: that of the digestive tract. It will be of interest to explore the cellular mechanisms underlying their differential sensitivity. These might result from differences in receptor levels or composition—the Ret-like receptor tyrosine kinase Stitcher, recently shown to synergize with InR in mitotic tissues, is a possible candidate (O’Farrell et al., 2013).

Figure 6. Distinct Effects on Energy Homeostasis Resulting from Pan-Tracheal or Gut-Specific Reductions in Tracheal Terminal Branching
(A) Reduced growth of most tracheal terminal cells (achieved using DSRF>btl-RNAi) does not affect the time between egg laying and pupation (only the two controls are significantly different from one another, p < 0.001, n = 40 larvae/set).
(B) This genetic manipulation leads to shorter-lived adult male flies in the presence of nutritious food (p < 0.0001 for all three comparisons, n = 70–120 flies/set).
(C) DSRF>btl-RNAi larvae have an increased length to width ratio (p < 0.001 versus GAL4 control, p < 0.0001 versus UAS control, n = 30 samples/set, total 300 larvae/set).
(D) They also have a reduced fat/protein content ratio (p < 0.0001 versus GAL4 control and p = 0.013 versus UAS control, n = 19 samples/set, total 190 larvae/set).
(E) An increase in free glycerol is also apparent in their hemolymph (p = 0.002 versus GAL4 control, p < 0.001 versus UAS control, n = 13 samples/set, total 130 larvae/set).
(F) A gut-specific reduction in tracheal terminal cell growth (achieved using DSRF>InR-RNAi) does not affect the survival of adult male flies in well-fed conditions (n = 60–140 flies/set).
(G) The same genetic manipulation leads to enhanced survival when adult male flies are subject to nutrient restriction (p < 0.0001 versus either control, p < 0.001 GAL4 versus UAS controls, n = 110–120 flies/set).
(H and I) The lipid stores of these adult males are relatively normal in well-fed conditions (H, p = 0.001 versus UAS control but not significant versus GAL4 control, p = 0.002 GAL4 versus UAS controls, n = 7 samples/set, total 70 flies/set), but they are more reduced than those of controls upon nutrient restriction (I, p = 0.002 versus either UAS or GAL4 controls, n = 7 samples/set, total 70 flies/set). See also Figure S7.
Alternatively, it could be caused by differences in downstream signaling components such as Foxo, which has been shown to account for some organ-specific responses (Tang et al., 2011).

Functionally, by uncovering gut-specific effects of tracheation on adult survival and lipid mobilization upon nutrient scarcity, we have identified the tracheal system as a possible anatomical substrate for the previously reported effects of nutrient acquisition during developmental and growth periods on a variety of adult features (Foley and Luckinbill, 2001; Zwaan et al., 1991). Enterocytes would appear to be the obvious cellular mediators of these effects; changes in oxygen supply may modulate the metabolic state of these absorptive cells, and long-term adaptations to nutrient scarcity may result from differential nutrient absorption and/or utilization. However, enterocytes need not be the only intestinal targets of the nutrient-driven tracheal changes: the tracheal regulation of stem cell proliferation described above (Li et al., 2013) provides an alternative (or additional) target. Consistent with this idea, there is correlogram as well as (more limited) functional data implicating neuronal factors in the regulation of angiogenesis in tumor environments (Jang et al., 2000; Madden et al., 2011; Toda et al., 2008). Furthermore, oxygen need not be the sole mediator of the gut tracheal-driven adaptations; Li et al. (2013) also found that tracheae produce Dpp, an important TGFβ-like signaling molecule. In future, it will be of interest to explore not only these intestinal targets, but also whether the intestinal tracheal plasticity is more widely regulated by other environmental stimuli—such as gut epithelial infection or damage. From a more translational perspective, most studies of adaptive angiogenesis in vertebrates have focused on the adipose vasculature (Cao, 2010; Lijnen, 2008). In light of our Drosophila findings, it will be of interest to explore the nutritional plasticity of the gastrointestinal vasculature, as well as its contribution to pathologies such as obesity or to the metabolic improvements following gastric bypass interventions.

**EXPERIMENTAL PROCEDURES**

**Visualization and Scoring of Tracheal Growth**

Tracheae were imaged and blindly scored using DIC optics (see Extended Experimental Procedures for details). Quantifications were performed as follows:

**Body Wall Tracheae**

The stereotypical endings of the third dorsal branch, directly posterior to the large tracheal commissure on the third segment, were counted as described in (Centanin et al., 2008).

**CNS Tracheae**

Tracheal coverage was quantified in the VNC—a relatively flat tissue with well-defined anatomical boundaries—as the ratio between the total length of tracheal arbours (which are complex and nonstereotypical) divided by total VNC area (μm²/μm²). Tracheal length was measured using a custom-written ImageJ macro (Schneider et al., 2012). After median filtering (radius = 3 pixels) to reduce image noise, a polygonal region of interest (ROI) was manually drawn to mark the tissue area. Following background subtraction to enhance the visibility of tracheae, the image was segmented and the tracheal area within the ROI was measured.

**Gut Tracheae**

In the mid-hindgut, where the tissue surface and three-dimensional properties allowed semiautomated quantification, the same procedure as for the VNC was used, but the segmented image was subsequently skeletonized. Parts of gut tissue wrongly identified as tracheae or segments of the tracheal tree missed by the program were manually edited before counting the total number of pixels in the skeletonized tracheal tree. In other intestinal portions, where the ruggedness and/or bends and twists of the target tissue made semiautomated quantification impractical, tracheal coverage was blindly scored using Likert-type scales ranging from no difference to wild-type (3) to strongly increased (5) or strongly reduced (1) (see Figure 1 legend for color coding of displays). The validity of this scoring system was confirmed in the body wall and mid-hindgut, where Likert-quantified scores were comparable to those obtained by counting or by semiautomated quantification respectively (data not shown). Likert rank data were displayed as the mean (circled) on diverging stacked bar charts, with the percentage of samples assigned to each Likert rank reflected in the length of each differently colored segment.

We refer to Extended Experimental Procedures for details of statistical analyses, fly stocks, diets, and more standard methods (immunohistochemistry, transmission electron microscopy, metabolic assays, survival assays, developmental rate and size quantifications, and in vivo recordings of neuronal activity).

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Extended Experimental Procedures, seven figures, and two movie and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2013.12.008.

**ACKNOWLEDGMENTS**

We thank Jordi Casanova, Jaya Krishnan, Stefan Luschnig, Helen Skaer, Pablo Wapnner, and members of the lab (Bhavana Chanana, Paola Cognigni, Esmeralda Parra-Peralbo, and Clara Recasens-Zorzo) for discussions and/or critical reading of the manuscript. We also thank Zoltán Soltész for the image analysis software used to analyse the GCaMP fluorescence. We are grateful to Li Liu and Yijin Wang for sharing results prior to publication and to Boris Adryan, Jordi Casanova, Louise Couton, Sebastian Groenke, Linda Partridge, Stefan Pulver, Orie Shafer, Maarten Zwart, the VDRC, and Bloomington Stock Center for providing fly stocks and antibodies. This work was funded by a Wellcome Trust RCDF to I.M.-A. (WT083559MA), a Wellcome Trust PhD studentship to G.A.L. (WT086807MA), an EMBO LTF to B.H., an ERC Starting Grant, a BBSRC project grant (BB/J007110/1) and the MRC. I.M.-A. is also supported by the EMBO Young Investigator Programme.

Received: May 13, 2013
Revised: August 26, 2013
Accepted: November 5, 2013
Published: January 16, 2014

**REFERENCES**

Adolphé, J.L., Whiting, S.J., Juurlink, B.H., Thorpe, L.U., and Alcorn, J. (2010). Health effects with consumption of the flax lignan secoisolariciresinol diglucoside. Br. J. Nutr. 103, 929–938.

Andersen, D.S., Colombani, J., and Léopold, P. (2013). Coordination of organ growth: principles and outstanding questions from the world of insects. Trends Cell Biol. 23, 336–344.

Asano, A., Morimatsu, M., Nikami, H., Yoshida, T., and Saito, M. (1997). Adrenergic activation of vascular endothelial growth factor mRNA expression in rat brown adipose tissue: implication in cold-induced angiogenesis. Biochem. J. 329, 179–183.

Belkacemi, L., Nelson, D.M., Desai, M., and Ross, M.G. (2010). Maternal undernutrition influences placental-fetal development. Biol. Reprod. 83, 325–331.

Böhni, R., Riesgo-Escovar, J., Oldham, S., Brogiolo, W., Stocker, H., Andruss, B.F., Beckingham, K., and Hafen, E. (1999). Autonomous control of cell and organ size by CHICO, a Drosophila homolog of vertebrate IRS1-4. Cell 97, 865–875.

---

**Cell** 156, 69–83, January 16, 2014 ©2014 The Authors 81
Brogiole, W., Stocker, H., Ikeya, T., Rintelen, F., Fernandez, R., and Hafen, E. (2001). An evolutionarily conserved function of the Drosophila insulin receptor and insulin-like peptides in growth control. Curr. Biol. 11, 213–221.

Cao, Y. (2007). Angiogenesis modulates adipogenesis and obesity. J. Clin. Invest. 117, 2362–2368.

Cao, Y. (2010). Adipose tissue angiogenesis as a therapeutic target for obesity and metabolic diseases. Nat. Rev. Drug Discov. 9, 107–115.

Carmeliet, P., and Tessier-Lavigne, M. (2005). Common mechanisms of nerve and blood vessel wiring. Nature 436, 193–200.

Centanin, L., Dekany, A., Romero, N., Irisari, M., Gorr, T.A., and Wappner, P. (2008). Cell autonomy of HIF effects in Drosophila: tracheal cells sense hypoxia and induce terminal branch sprouting. Dev. Cell 14, 547–558.

Chaudhuri, A., Dandona, P., and Fonseca, V. (2012). Cardiovascular benefits of exogenous insulin. J. Clin. Endocrinol. Metab. 97, 3079–3091.

Cheili, J.M., and Brand, A.H. (2010). Nutrition-responsive glia control exit of neural stem cells from quiescence. Cell 143, 1116–1173.

Cheung, U., Moghaddasi, M., Hall, H.L., Smith, J.J., Buck, L.T., and Woodin, S.K., and Singh, B. (2013). Probiotic metabolites as epigenetic targets in the prevention of colon cancer. Nutr. Res. 33, 182–196.

Chet, R.J., Klein, O.D., Martin, G.R., and Krasnow, M.A. (2008). The branching programme of mouse lung development. Nature 453, 745–750.

Miguel-Alia, I., Thor, S., and Gould, A.P. (2008). Postmitotic specification of Drosophila insulinergic neurons from pioneer neurons. PLoS Biol. 6, e58.

Mukouyama, Y.S., Shin, D., Britsch, S., Taniguchi, M., and Anderson, D.J. (2002). Sensory nerves determine the pattern of arterial differentiation and blood vessel branching in the skin. Cell 109, 693–705.

Mukouyama, Y.S., Gerber, H.P., Ferrara, N., Gu, C., and Anderson, D.J. (2005). Peripheral nerve-derived VEGF promotes arterial differentiation via neuropilin 1-mediated positive feedback. Development 132, 941–952.

O’Brien, L.E., Soliman, S.S., Li, X., and Bilder, D. (2011). Altered modes of stem cell division drive adaptive intestinal growth. Cell 147, 603–614.

O’Farrell, F., Wang, S., Katheder, N., Rusten, T.E., and Samakovlis, C. (2013). Two-tiered control of epithelial growth and autophagy by the insulin receptor and the ret-like receptor, stitcher. PLoS Biol. 11, e1001612.

Ogasawara, M., Murata, J., Kamitani, Y., Hayashi, K., and Saiki, I. (1999). Inhibition by vasoactive intestinal polypeptide (VIP) of angiogenesis induced by murine Colon 26-L5 carcinoma cells metastasized in liver. Clin. Exp. Metastasis 17, 283-291.

Pardal, R., and López-Barneo, J. (2002). Low glucose-sensing cells in the carotid body. Nat. Neurosci. 5, 197–198.

Potente, M., Gerhardt, H., and Carmeliet, P. (2011). Basic and therapeutic aspects of angiogenesis. Cell 146, 873–887.

Prabakar, N.R. (2000). Oxygen sensing by the carotid body chemoreceptors. J. Appl. Physiol. 88, 2287–2295.

Quaegebeur, A., Lange, C., and Carmeliet, P. (2011). The neurovascular link in health and disease: molecular mechanisms and therapeutic implications. Neuron 71, 406–424.

Ribatti, D., Conconi, M.T., and Nussdorfer, G.G. (2007). Nonclassic endogenous novel [corrected] regulators of angiogenesis. Pharmacol. Rev. 59, 185–205.

Rullifson, E.J., Kim, S.K., and Nusse, R. (2002). Ablation of insulin-producing neurons in flies: growth and diabetic phenotypes. Science 296, 1118–1120.

Rutland, C.S., Latunde-Dada, A.O., Thorpe, A., Plant, R., Langley-Evans, S., and Leach, L. (2007). Effect of gestational nutrition on vascular integrity in the murine placenta. Placenta 28, 734–742.

Schneider, C.A., Rasband, W.S., and Eliceiri, K.W. (2012). NIH Image to ImageJ: 25 years of image analysis. Nat. Methods 9, 671–675.

Slaiby, M., Delanoue, R., Gronke, S., Prabaker, N.R., and Léopold, P. (2009). A Drosophila insulin-like peptide promotes growth during nonfeeding states. Dev. Cell 17, 874–884.

Sousa-Nunes, R., Yee, L.L., and Gould, A.P. (2011). Fat cells reactivate quiescent neuroblasts via TOR and glial insulin relays in Drosophila. Nature 471, 508–512.

Storkebaum, E., Quaegebeur, A., Vikkula, M., and Carmeliet, P. (2011). Cerebrovascular disorders: molecular insights and therapeutic opportunities. Nat. Neurosci. 14, 1390–1397.

Sun, K., Wernstedt Astholm, I., Kusminski, C.M., Bueno, A.C., Wang, Z.V., Pollard, J.W., Breken, R.A., and Scherer, P.E. (2012). Dichotomous effects of VEGF-A on adipose tissue dysfunction. Proc. Natl. Acad. Sci. USA 109, 5874–5879.
Sung, H.K., Doh, K.O., Son, J.E., Park, J.G., Bae, Y., Choi, S., Nelson, S.M., Cowling, R., Nagy, K., Michael, I.P., et al. (2013). Adipose vascular endothelial growth factor regulates metabolic homeostasis through angiogenesis. Cell Metab. 17, 61–72.

Suska, A., Miguel-Aliaga, I., and Thor, S. (2011). Segment-specific generation of Drosophila Capability neuropeptide neurons by multi-faceted Hox cues. Dev. Biol. 353, 72–80.

Taghert, P.H., and Nitabach, M.N. (2012). Peptide neuromodulation in invertebrate model systems. Neuron 76, 82–97.

Talsma, A.D., Christov, C.P., Terriente-Felix, A., Linneweber, G.A., Perea, D., Wayland, M., Shafer, O.T., and Miguel-Aliaga, I. (2012). Remote control of renal physiology by the intestinal neuropeptide pigment-dispersing factor in Drosophila. Proc. Natl. Acad. Sci. USA 109, 12177–12182.

Tang, H.Y., Smith-Caldas, M.S., Driscoll, M.V., Salhadar, S., and Shingleton, A.W. (2011). FOXO regulates organ-specific phenotypic plasticity in Drosophila. PLoS Genet. 7, e1002373.

Tian, L., Hires, S.A., Mao, T., Huber, D., Chiappe, M.E., Chalasani, S.H., Petreanu, L., Akerboom, J., McKinney, S.A., Schreiter, E.R., et al. (2009). Imaging neural activity in worms, flies and mice with improved GCaMP calcium indicators. Nat. Methods 6, 875–881.

Toda, M., Suzuki, T., Hosono, K., Hayashi, I., Hashiba, S., Onuma, Y., Amano, H., Kurihara, Y., Kurihara, H., Okamoto, H., et al. (2008). Neuronal system-dependent facilitation of tumor angiogenesis and tumor growth by calcitonin gene-related peptide. Proc. Natl. Acad. Sci. USA 105, 13550–13555.

Tonello, C., Giordano, A., Cozzi, V., Cinti, S., Stock, M.J., Carruba, M.O., and Nisoli, E. (1999). Role of sympathetic activity in controlling the expression of vascular endothelial growth factor in brown fat cells of lean and genetically obese rats. FEBS Lett. 442, 167–172.

Uv, A., Cantera, R., and Samakovlis, C. (2003). Drosophila tracheal morphogenesis: intricate cellular solutions to basic plumbing problems. Trends Cell Biol. 13, 301–309.

Vesalius, A. (1543). De humani corporis fabrica (Basel: Oporinus).

Willecke, M., Toggweiler, J., and Basler, K. (2011). Loss of PI3K blocks cell-cycle progression in a Drosophila tumor model. Oncogene 30, 4067–4074.

Zwaan, B.J., Bijlsma, R., and Hoekstra, R.F. (1991). On the developmental theory of ageing. I. starvation resistance and longevity in Drosophila melanogaster in relation to pre-adult breeding conditions. Heredity (Edinb) 66, 29–39.