The Polycomb Repressive Complex 2.1 Links the Food Environment to a Persistent Neural State

Anoumid Vaziri¹,², Morteza Khabiri³†, Brendan T. Genaw⁵, Christina E. May²,⁶, Peter L. Freddolino³,⁴, Monica Dus¹,²,⁵,⁶*

¹The Molecular, Cellular and Developmental Biology Graduate Program, The University of Michigan, Ann Arbor, MI 49109
²Department of Molecular, Cellular and Developmental Biology, College of Literature, Science, and the Arts, The University of Michigan, Ann Arbor, MI 49109
³Department of Biological Chemistry, The University of Michigan, Ann Arbor, MI 48109, USA
⁴Department of Computational Medicine and Bioinformatics, The University of Michigan, Ann Arbor, MI 48109, USA
⁵Program in Biology, College of Literature, Science, and the Arts, The University of Michigan, Ann Arbor, MI, 48109, USA
⁶The Neuroscience Graduate Program, The University of Michigan, Ann Arbor, MI 49109
†Current address: Department of Biological Sciences, Quinnipiac University, Hamden, CT 06518
*
* Correspondence: mdus@umich.edu
Abstract

Interactions between genes and environment sculpt the responses of cells to environmental stimuli. In neuronal cells this process can lead to long term changes in the behavioral repertoire of animals, which in turn impacts disease risk. Here we show that the Polycomb Repressive Complex 2.1 (PRC2.1) modulates the physiology of sweet gustatory neurons and the taste behavior of *D. melanogaster* fruit flies in response to the food environment. A high sugar diet caused a redistribution of PRC2.1 chromatin occupancy resulting in the repression of a transcriptional network required for the responsiveness of the gustatory neurons to sweet stimuli. These changes led to lower sweet sensation, which in turn promoted obesity. Nearly half of the transcriptional changes mediated by PRC2.1 on a sugar diet persisted when animals were moved back to a control diet, causing a permanent decrease in sweet taste that was dependent on the constitutive activity of PRC2.1. Thus, our results point to a novel mechanism involved in modulating neural plasticity, behavior, and disease in response to the food environment.
Introduction

In 1958 D. L Nanney coined the term “epigenetic control system” to explain phenotypic variation that was not dependent on the “genetic library” (Nanney, 1958). More than half a century later, we understand that epigenetic mechanisms sculpt cellular responses to environmental variation via changes in gene expression (Allis and Jenuwein, 2016). These responses show varying degrees of stability, and some phenotypes persist even after the environmental signals disappear, forming a memory of past experiences. This process has far-reaching implications for all cells, but it is particularly critical for neurons, because their molecular properties direct the physiology and behavior of the whole organism. Indeed, environmental conditions that permanently alter neurons reshape the way in which future stimuli are perceived and processed, limiting the behavioral repertoire of animals and influencing disease risk (Campbell and Wood, 2019; Dulac, 2010; Sweatt, 2013). The effects of environmental reprogramming on behavioral plasticity have been elegantly studied in the context of maternal care, learning and memory, and addiction (Champagne, 2008; Dulac, 2010; Nestler and Lüscher, 2019; Sweatt, 2013), which has led to the identification of some of the molecular mechanisms of anxiety, depression, and cognitive ability (Campbell and Wood, 2019). However, because these studies were done in whole brain regions such as the hippocampus and striatum, we still do not know how environmental signals impact the function of specific neural circuits to persistently reprogram behavior. Further, the mechanisms through which other biomedically important environments – such as diet and pollutants – alter brain physiology and behavior remain largely undefined. Here we studied the effects of high dietary sugar on taste sensation as a model to identify the molecular mechanisms through which the food environment reprograms neural physiology and behavior.
Like many behaviors, the ability of animals to detect food sources is under the control of both genes and environment. The genome shapes the sensory apparatus of each species to match their unique ecological niche, but despite this genetic constraint, this system is plastic and can be altered by diet composition (Beauchamp and Jiang, 2015). Indeed, diet has been shown to change the taste perceptions of animals from insects to humans (Bertino et al., 1982; Glendinning et al., 2001; May et al., 2019a; Stewart and Keast, 2012; Wang et al., 2016; Wise et al., 2016; Zhang et al., 2013), and across organisms, sensory alterations affect food preference and intake, influencing the development of conditions like obesity and heart disease (Bertino et al., 1982; May et al., 2019a; Stewart and Keast, 2012). We recently found that high dietary sugar dulls the responses of the fruit fly taste neurons to sweet stimuli, causing higher food intake and weight gain (May et al., 2019a, 2019b). Mammals fed high nutrient diets also show changes in taste, neural responses, and food preferences (Ahart et al., 2019; Chen et al., 2010; Crow, 2012; Kaufman et al., 2018; Maliphol et al., 2013; May et al., 2019a; McCluskey et al., 2020; Sartor et al., 2011; Weiss et al., 2019), arguing that the effects of diet on taste are conserved and promote obesity. However, the molecular mechanisms through which the food environment alters taste sensation, and more broadly, neural physiology, are largely unknown. Here we exploited the exquisite genetics tools of the fruit fly and the relative simplicity of its sensory system to tackle these questions. We found that the chromatin silencing Polycomb Repressive Complex 2.1 (PRC2.1) tuned the activity of the sweet sensory neurons and taste sensation to the food environment by repressing a transcriptional program that shapes the synaptic, signaling, and metabolic properties of these cells. Interestingly, this diet-dependent transcriptional remodeling persisted even when animals were returned to the control diet, leading to lasting changes in sweet taste sensation that depended on the constitutive activity of PRC2.1. Together our findings suggest that the food environment reprograms sensory
responses by activating epigenetic mechanisms that persistently restrict the perception of future stimuli, alter behavior, and increase the risk for obesity and metabolic disease.

Results

PRC2.1 modulates sweet taste in response to the food environment

*Drosophila melanogaster* fruit flies fed high dietary sugar experience lower sweet taste sensation as a result of the decreased responsiveness of the sweet sensory neurons to sugar stimuli (May et al., 2019a). Given the importance of sensory cues to control eating, and recent data that diet also impacts taste in mammals (Ahart et al., 2019; Chen et al., 2010; Kaufman et al., 2018; Maliphol et al., 2013; May et al., 2019a; McCluskey et al., 2020; Weiss et al., 2019), we set out to identify the molecular mechanisms through which the food environment shapes sensory responses. Since sweet taste deficits develop within 2-3 days upon exposure to the high sugar diet and independently of weight gain (May et al., 2019a), we reasoned that gene regulatory mechanisms may be involved in modulating the responses of the sensory neurons. To test this hypothesis we conducted a screen for gene regulatory factors necessary for sweet taste defects on a high sugar diet. To do this, we fed control (w1118cs) and mutant flies a control diet (CD, ~5% sucrose) or a diet supplemented with 30% sucrose (sugar diet, SD) for 7 days and then tested their ability to taste using the Proboscis Extension Response (Shiraiwa and Carlson, 2007). This behavioral assay measures taste responses by quantifying the amount of proboscis extension (0= no extension, 0.5=partial extension, 1= full extension) when the fly labellum – where the dendrites and cell bodies of the taste neurons are located (Fig. S1A) – is stimulated with three different concentrations of sucrose (30%, 10%, 5%); this generates a taste curve where flies respond more intensely to higher sugar stimuli (Fig. 1B, gray circles). Flies fed a sugar diet show a marked decrease in PER compared to control diet flies (Fig. 1B, gray
squares); however, mutants for the core Polycomb Repressive Complex 2 (PRC2) – which
includes the histone 3 lysine 27 (H3K27) methyltransferase Enhancer of Zeste (E(z)), and the
obligate accessory factors Suppressor of zeste 12, (Su(z)12) and extra sex combs, (esc) (Fig. 1A) – had largely the same proboscis extension response (PER) on a control and sugar diet
(Fig. 1B, right, red shades). To confirm the role of PRC2 in taste sensation, we supplemented
the control and sugar diet with EED226, a PRC2 inhibitor (herein referred to as EEDi) that
destabilizes the core complex by binding to the tri-methyl H3K27 (H3K27me3) binding pocket of
EED (the homologue of esc in M. musculus) (Qi et al., 2017). While animals fed a sugar diet
plus vehicle (10% DMSO) experienced lower PER, those fed a SD+EEDi retained normal sweet
taste responses (Fig. 1C), consistent with results from the PRC2 mutants. Thus, mutations and
inhibition of PRC2 rescue the blunting of sweet taste that occurs in the high sugar food
environment.

In flies PRC2 forms two main subcomplexes, PRC2.1 and PRC2.2, which contain
distinct accessory factors that influence the targeting of the core complex to the genome
(Laugesen et al., 2019). Mutations in the Polycomb-like (Pcl) gene, the accessory factor to
PRC2.1, phenocopied PRC2 mutants and rescued sweet taste deficits in flies fed a sugar diet
(Fig. 1D). In contrast, flies with deficits in the PRC2.2-members Jumonji, AT rich interactive
domain 2 (Jarid2) and jing still showed a blunting of sweet taste responses in flies fed a sugar
diet (Fig. S1B). Interestingly, members of the Polycomb Repressive Complex 1 (PRC1) and the
recruiter complex PhoRC were also not required for taste changes in responses to a sugar diet
(Fig. S1C and Fig. S1D-E). Thus, the PRC2.1 complex is necessary for the sensory changes
that occur in the high sugar environment.

We next asked if PRC2.1 is required specifically in the sweet sensory neurons to
decrease their responses to sweet stimuli on the sugar diet. To do this, we used the GAL4/UAS
system to knock down Pcl in the sweet taste neurons using the *Gustatory receptor 5a GAL4* driver, Gr5a-GAL4, which labels ~60 cells in the proboscis of adult flies (Chyb et al., 2003); we selected Pcl to narrow the effect to the PRC2.1 complex. Knockdown of Pcl in Gr5a+ neurons using two independent RNAi transgenes (50% knockdown efficiency, Fig. S2A) prevented sweet taste deficits in animals fed a sugar diet (Fig. 1E, and Fig. S2B). Pcl knockdown, however, had no effect on a control diet (Fig. S2C), in accordance with the observation that E(z) and Pcl mutants have no effect on taste on a control diet (Fig. 1B, C) and suggesting that these phenotypes are uncovered only by the high sugar food environment.

Since Pcl is thought to target the PRC2 core complex to chromatin (Laugesen et al., 2019), we hypothesized that its overexpression may be sufficient to induce sweet taste deficits even in the absence of a high sugar food environment. Indeed, overexpression of Pcl specifically in the Gr5a+ neurons induced sweet taste deficits in flies fed a control diet compared to transgenic controls (Fig. 1F). The effects of Pcl overexpression were abolished by treatment with the PRC2 inhibitor EEDi (Fig. 1G), arguing that Pcl causes sweet taste deficits entirely through the action of PRC2 and not through some yet unidentified mechanism. Importantly, Pcl overexpression had no effect on the number of Gr5a+ neurons in the proboscis (Fig. S2D), and so the taste deficits cannot be attributed to a decrease in the number of cells. To exclude the possibility that the effects of manipulating Pcl were developmental, we used the temperature sensitive *tubulin-GAL80* transgene to limit expression of *UAS-Pcl* and *PclRNAi* only in adult flies. Switching the flies to the non-permissive temperature and diet 4 days post eclosion, resulted in the same effects on sweet taste as using the Gr5a-GAL4 alone (Fig. S2E). Together, these experiments establish that PRC2.1 is required cell-autonomously in the Gr5a+ neurons to mediate the effects of a high sugar diet on sweet taste.
**Pcl mutant flies have normal sensory responses and are resistant to diet-induced obesity**

Flies on a high sugar diet have lower sweet taste because the neural responses of the taste neurons to sweet stimuli are decreased (May et al., 2019a). Since Pcl mutants have identical taste on a control and sugar diet, we hypothesized that the responses of the sensory neurons to sucrose stimulation should also be similar. To test this, we expressed the genetically encoded presynaptic calcium indicator UAS-GCaMP6s-Brp-mCherry (Kiragasi et al., 2017) in the sweet sensing neurons and measured their *in vivo* responses to stimulation of the proboscis with 20% sucrose in Pcl mutant animals (Fig. 2A). Indeed, the responses to sucrose stimulation were identical in Pcl mutant flies fed a control diet and sugar diet (Fig. 2B), matching the behavioral data (Fig. 1); importantly, this rescue was not due to an increase in the number of sweet taste cells (Fig. 2C).

We previously showed that restoration of sweet taste neuron activity in flies fed high dietary sugar protected them from diet-induced obesity (May et al., 2019a). Since Pcl mutants abolished the deficits in neural and behavioral responses to sweetness in animals fed a high sugar diet, we anticipated that they should also prevent a diet-dependent increase in triglycerides. Indeed, sugar-diet flies with knockdown of Pcl in the Gr5a+ neurons remained as lean as animals on a control diet (Fig. 2D), while triglycerides increased in control flies fed a sugar diet (Fig. 2D). Importantly, there was no difference in the levels of triglycerides between control and Pcl knockdown flies fed a control diet (Fig. 2D). Together, these data suggest that, in response to the food environment, Pcl modulates the responsiveness of the sweet gustatory neurons to promote diet-induced obesity.

**Pcl chromatin occupancy is redistributed in the high sugar environment**
Our experiments show that PRC2.1 plays a critical role in the neural activity, behavior, and the metabolic state of animals exposed to the high sugar food environment. To identify the molecular mechanisms underlying these phenotypes, we measured the chromatin occupancy of \textit{Pcl} in the \textasciitilde60 \textit{Gr5a}+ neurons using Targeted DNA Adenine Methyltransferase Identification (TaDa) (van Steensel and Henikoff, 2000) (Southall et al., 2013) (Marshall et al., 2016). To do this we generated \textit{Dam::Pcl} (UAS-LT3-Dam::Pcl) transgenic flies and compared them to \textit{Dam-only} flies (UAS-LT3-Dam) to control for non-specific methylation by the freely diffusing \textit{Dam} protein (van Steensel and Henikoff, 2000) (Fig. 3A) and obtain a measure of chromatin accessibility \textit{in vivo} (CATaDa) (Aughey et al., 2018). To specifically profile \textit{Pcl} binding to chromatin in the sweet sensory neurons and limit the induction of \textit{Dam}, we expressed the \textit{Dam::Pcl} and \textit{Dam} transgenes in combination with \textit{Gr5a-GAL4; tubulin-GAL80\textasciitilde}. To induce the expression of each UAS transgene we shifted the flies to the permissive temperature (28°C) for 20 hours after they had been exposed to a control or sugar diet for 3 days (Fig. 3A). We selected this time point because we previously showed that sweet taste defects developed within 3 days of exposure to the sugar diet (May et al., 2019a).

Most of the variation in the biological replicates of \textit{Dam::Pcl} normalized to \textit{Dam} alone (see Methods) was due to diet (Fig. S3A), consistent with the high Pearson correlations within each dietary condition (Fig. S3B). Further, the accessibility profile of \textit{Dam} at the \textit{Gr5a} sweet taste receptor gene promoter was high, while that at the \textit{Gr66a} bitter taste receptor promoter—which is only expressed in bitter cells, closely located near the sweet cells—was low (Fig. 3B), suggesting that the transgenes were appropriately targeted to the sweet taste neurons and that the limited induction controlled for background DNA methylation.

We first analyzed \textit{Pcl} chromatin occupancy in the \textit{Gr5a}+ neurons of flies on a control diet by comparing our data to a previous study that annotated five major chromatin types in \textit{D}. 
melanogaster using a similar technique (DNA Adenine Methyltransferase Identification, Dam-ID) (Filion et al., 2010). Pcl targets were enriched in Polycomb chromatin (blue), compared to other chromatin types (green and black = repressive; red and yellow= active) (Fig. 3C); for example, Pcl occupancy was high and Dam accessibility low at two known Polycomb blue chromatin clusters, (Fig. S3E), while the opposite was true for regions in other chromatin types (Fig. S3F, red and yellow) (Filion et al., 2010). We next asked whether Pcl was enriched at Polycomb Response Elements (PREs), cis-regulatory sequences to which Polycomb Group Proteins bind in D. melanogaster (Chan et al., 1994) (Müller and Kassis, 2006). Using a recently developed tool that predicts PRE regions genome-wide (Khabiri and Freddolino, 2019), we found that Pcl was present in these regions (Fig. 3D, gray line), with an enrichment for intergenic (3.2-fold enriched, p<0.001, Monte Carlo permutation test) and enhancer PREs (2.9-fold enriched, p<0.001, Monte Carlo permutation test).

To determine changes in Pcl occupancy with diet, we compared Pcl peaks between flies fed a control and sugar diet. While ~70% of the Pcl peaks were shared between the control (CD) and sugar diet (SD) (Fig. S3C, D), we found more Pcl at PREs on a sugar diet compared to a control diet (Fig. 3D, purple line). Interestingly, chromatin accessibility at both Pcl peaks (Fig. 3E) and PREs (Fig. 3F) was decreased in the sugar diet condition. Our analysis also showed that differentially bound peaks had a 3.3-fold enrichment of overlap for enhancer-type PREs (p<0.001, Monte Carlo permutation test). Further examination of the differentially bound genes, revealed a redistribution in Pcl occupancy (Fig. S3G), with a similar number of genes with higher (group 1) and lower (group 2) Pcl binding on a sugar diet compared to the control diet (Fig. 3G). Using iPAGE for pathway enrichment analysis (Goodarzi et al., 2009) we found that most of the genes differentially bound by Pcl were transcription factors with both promoter and enhancer binding. Notably, transcription factors implicated in axon target recognition and
nervous system development showed an enrichment in \( Pcl \) binding on a sugar diet, while those involved in proboscis development and feeding behavior had both an increase and decrease in \( Pcl \) occupancy on a sugar diet (Fig. 3H, for full iPAGE see Fig. S4). While the large majority of genes differentially bound by \( Pcl \) were in the gene regulation category (80%), the pathway enrichment analysis also uncovered a few metabolism Gene Ontology (GO) terms (Fig. 3H and Fig. S4). In summary, we found that PRC2.1 targeted transcription factors involved in neuronal processes and development in the \( Gr5a^+ \) neurons, and its chromatin occupancy was redistributed at these loci in the high sugar diet environment. This redistribution could result in changes in the expression of these transcription factors and their targets, and in turn, affect the sensitivity of the sensory neurons and taste sensation.

**PRC2.1 sculpts the transcriptional responses of the \( Gr5a^+ \) neurons in response to diet**

To test the hypothesis that redistribution of PRC2.1 chromatin occupancy alters the physiology of the sweet sensing neurons by remodeling transcription, we used Translating mRNA Affinity Purification (TRAP) (Chen and Dickman, 2017)) to isolate mRNAs associated with the ribosomes of these cells. (Fig. 4A). To capture the dynamics of this process, we collected samples from age-matched \( Gr5a^{+}\text{Rpl3-3XFLAG} \) flies fed a sugar diet for 3 and 7 days (Fig. S5A). We first verified that this technique selected for mRNAs in the \( Gr5a^+ \) neurons alone by quantifying the normalized read counts (\( Gr5a^+ / \text{input} \)) for three sweet taste receptor genes \( Gr5a, Gr64f, \) and \( Gr64a, \) which are expressed in cells labeled by the \( Gr5a\)-GAL4. Indeed, these transcripts were enriched in the \( Gr5a^+ \) fraction compared to the input (Fig. S5B), while the opposite was true for the bitter receptor gene \( Gr66a, \) which is expressed only in the bitter sensing neurons the taste sensilla (Scott, 2018) (Fig. S5B).
Notably, we observed a large negative skew in gene expression in the Gr5a+ neurons of flies fed a sugar diet for 3 (SD3, mint; compared to the control diet) and 7 days (SD7, teal; compared to the control diet) (Fig. 4B-C, -0.8 and -1.7 skew), with ~68% and 87% of differentially expressed genes (DEG, each compared to control diet, Wald test, q< 0.1) showing negative log₂ fold changes (l2fc) respectively (Fig. 4B-C, SD3 and SD7). Overall, we found ~800 transcripts differentially expressed at each time point compared to the control diet, while ~190 changed at both time points (Fig. 4D, Venn diagram, Wald test, q< 0.1). Gene Ontology (GO) analysis (Goodarzi et al., 2009) revealed that these genes were part of biological pathways involved in 3 broad categories: neural function/signaling, metabolism, and gene regulation (Fig. S6 and Fig. S7). GO terms for neuron-specific processes, such as dendritic membrane, sensory perception of chemical stimulus, and presynaptic/vesicle transport, were enriched at both timepoints (Fig. S6 and Fig. S7), suggesting that a high sugar diet may alter the physiology of the sensory neurons through these pathways. Interestingly, flies fed a sugar diet for 7 days also showed changes in GO terms linked to neurodevelopmental processes, such as asymmetric neuroblast division and neuron projection morphogenesis, indicating that longer exposure to the diet led to additional alterations in neural function. GO terms associated with metabolic changes were present in higher numbers in flies fed a sugar diet for 7 days, consistent with the findings that longer exposure to the high sugar diet leads to higher fat accumulation (May et al., 2019a). Finally, we observed changes in “regulatory” GO terms such as transcription factor and corepressor, consistent with our chromatin binding experiments (TaDa). Thus, consumption of a high sugar diet altered neural, regulatory, and metabolic genes in the Gr5a+ cells.

To determine the role of PRC2.1 in these changes, we performed the transcriptional profiling experiments in the Gr5a+ neurons of Pcl<sup>+/429</sup> mutant animals fed a control diet and sugar diet for 7 days (CD and SD7) (Fig. S5C). Strikingly, the Pcl mutation abolished the negative
skew (Fig. S5D) and largely nullified the effects of the high sugar diet environment on gene expression. Specifically, of the genes repressed by a sugar diet (Fig. 4D, heatmap) 32% had a positive log₂ fold change (Wald test, \( q < 0.1 \)) and 76% were unchanged (\( q < 0.1 \), practical equivalence test using a null hypothesis of a change of at least 1.5-fold; see Methods for details) between \( Pcl \) mutants fed a control and sugar diet. This effect was reflected in the GO analysis where terms changed by a high sugar diet in \( Pcl \) wild-type animals, such as dendritic membrane, sensory perception of chemical stimuli, synapse, and carbohydrate metabolic process, showed opposite trends in log₂ fold changes in \( Pcl \) mutants (Fig. S8). Thus, \( Pcl \) mutations abolished nearly all the gene expression changes induced by a high sugar diet consistent with their effects on behavior (PER, Fig.1 ), neural function (\textit{in vivo} calcium imaging, Fig. 2), and metabolism (triglycerides, Fig. 2). Together, these findings support the hypothesis that PRC2.1 tunes taste sensation to the food environment by influencing the expression of genes involved in the physiology of the sensory neurons.

**PRC2.1 represses a transcriptional program required for sweet taste**

We discovered that a high sugar diet environment repressed gene expression in the sweet sensory neurons, and that \( Pcl \) mutations almost entirely abolished its effect. This, together with the discovery that \( Pcl \) binding primarily changed at the enhancers of transcription factor genes on a sugar diet, suggests that \( Pcl \) redistribution may affect the expression of transcription factors that, in turn, control genes responsible for the overall responsiveness of these sensory neurons to sweetness. This idea was supported by the observation that \( Pcl \)-bound genes had lower expression levels than those not bound by it in the Gr5a+ neurons (Fig. 4E), with many genes showing high binding and low expression (\( \log_{2} \text{tpm} < 2 \), dark purple), while others having higher RNA read counts (\( \log_{2} \text{tpm} > 5 \), light purple) (Fig. S5E). To further test
this hypothesis, we looked for transcription factors that were directly bound by Pcl and that showed changes in gene expression on a sugar diet (Fig. S5F). This analysis revealed 5 transcription factors: four of these were activators – GATAe (Zn finger), nubbin/pdm (nub, POU homeobox), Ptx1 (paired-domain homeobox), and caudal (cad, hox-like homeobox) – which had higher Pcl binding (Fig. 5A) and lower mRNA levels on a sugar diet (Fig. 4F). The fifth transcription factor was the suppressor scarecrow (Scro, NK-like homeobox), which had lower Pcl binding (Fig. 5A) and higher mRNAs levels on a sugar diet (Fig. 4F). Notably, mutations in Pcl reversed the effects of a high sugar diet on the expression of these 5 genes, suggesting that the binding of Pcl modulates their mRNA levels (Fig. 4F). Interestingly, cad, Ptx1, GATAe, nub were enriched in the Gr5a+ neurons compared to the input (heads), while Scro was depleted in these cells (Fig. S9A). To test the effects of these five transcription factors on sweet taste, we manipulated their expression to mimic the direction of change on a high sugar diet. Knockdown of cad, Ptx1, GATAe, or nub, and overexpression of Scro in the Gr5a+ neurons of flies fed a control diet led to a decrease in sweet taste sensation (Fig. 5B) comparable to that experienced by animals on a sugar diet (Fig. 1B). Thus, cad, Ptx1, GATAe, nub are direct targets of PRC2.1 and necessary for sweet taste, while overexpression of Scro is sufficient to decrease it. However, overexpression of each activator alone and knockdown of Scro was not sufficient to rescue sweet taste in flies fed a sugar diet (Fig. S9B).

Given that the 4 activators are required for sweet taste sensation, we reasoned that they may control the expression of genes important for the proper function of the Gr5a+ neurons and normal sweet taste. To identify candidate target genes, we tiled the entire fruit fly genome using the motifs for each of these 5 transcription factors, converted the hits for each transcription factor to robust z-scores, and determined candidate regulatory targets based on estimates of the z-score threshold for binding in each case (Fig. S10A; see Methods for details). We then
flagged as “targets” the set of genes that had a putative binding site (exceeding our transcription factors-specific z-score cutoff) within a 2 kb region upstream of the annotated ORF start (Fig. S10B-F), and examined their expression pattern in the Gr5a+ neurons of flies on a control and sugar diet. This analysis revealed 658 genes that were collectively regulated by these five transcription factors. Targets of the transcriptional activators Cad, Pttx1, GATAe and Nub-- which Pcl repressed on a sugar diet-- showed negative log2 fold changes on a sugar diet (Fig. 5C, SD7, teal) that reverted in Pcl mutants (Fig. 5C, pink). Conversely, targets of the transcriptional repressor Scro – which was released from Pcl binding and had higher mRNA levels on a sugar diet – showed negative log2 fold changes on a sugar diet (Fig. 5C, SD7, teal) that reverted in Pcl mutants (Fig. 5C, pink). Strikingly, the 658 putative targets of these 5 transcription factors accounted for nearly all the genes changed by a high sugar diet (Fig. 4), suggesting that by directly modulating the expression of Pttx1, cad, GATAe, nub, and Scro, PRC2.1 could influence the neural state of the Gr5a+ neurons via their targets. When we examined whether these targets were regulated by more than one of the 5 transcription factors targeted by Pcl, we found significant overlap among the regulons of all of the 4 transcriptional activators with the exception of Pttx1-nub ( Fisher's exact test, FDR-corrected p<0.000001) (Fig. 5D), suggesting that they could direct the expression of a common set of genes. Further, a portion of Scro targets also overlapped with the regulons of each of the activators (Fisher’s exact test, q<0.000001), indicating that Scro could further drive the negative skew in gene expression on a high sugar diet beyond that caused by the direct binding of PRC2.1 to the activators. Transcription factors that share common targets are often part of feed-forward loops, where they regulate one another and themselves to ensure stability of gene expression. Indeed we found that GATAe had binding sites in the promoters of all four regulators considered here (cad, Scro, Pttx1, nub), in addition to binding its own promoter in an auto-regulatory loop (Fig.
Furthermore, Cad also targeted itself, Nub was one of Ptx1 targets, and Scro regulated both Cad and GATAe, forming a negative feedback loop with the latter (Fig. 5D). Thus, the 5 transcription factors differentially bound by PRC2.1 on a high sugar diet, form a hub that seems to regulate the properties of the Gr5a+ neurons. To gain a deeper understanding of these properties, we used pathway enrichment analysis on the regulons for each transcription factor. GATAe targets, which comprise a large number of the genes regulated by the 4 other transcription factors, were enriched for GO terms involved in synaptic assembly and growth, terminal bouton, neural projection morphogenesis, and protein kinase regulation (summarized in Fig. 5D, and Fig. S11). In contrast, Ptx1 targets were enriched in GO terms implicated in cyclic AMP signaling, detection of chemical stimuli, and morphogenesis (summarized in Fig. 5D, and Fig. S13), cad targets showed enrichments in adenylate cyclase activity, sensory perception, and neuropeptide signaling (summarized in Fig. 5D, and Fig. S12), and nub targets in calcium signaling and nucleosome (summarized in Fig. 5D, and Fig. S13). The targets of the repressor Scro showed enrichment in both neural and metabolic GO terms such as olfactory behavior and carbohydrate metabolic process (summarized in Fig. 5D, and Fig. S12).

To test the possibility that these targets form a functional network, we used STRING (Szklarczyk et al., 2019) and found a significant number of edges above the expected number (protein-protein interaction enrichment of $p < 1.0e-16$) suggesting that the targets are indeed part of a functional and biologically connected network in the Gr5a+ neurons. We then used a subset of the neural targets to build a second network to identify candidate target genes that may play a direct role in neural physiology and sweet taste. This network showed strong interactions between genes involved in synaptic organization and signal transduction and their connection with the upstream regulators (Fig. S14A). We chose two genes at the edges of the network, which are less likely to have redundant functions, the Adenylyl Cyclase X D (ACXD)
gene (Ueno and Kidokoro, 2008) and the Activity Regulated Cytoskeleton Associated Protein 1 (Arc1) (Ashley et al., 2018), which are involved in the sensing of stimuli and in synaptic plasticity, respectively. Knockdown and mutations of Arc1 or ACXD in the Gr5a+ cells of flies on a control diet led to a significant decrease in sweet taste responses compared to the transgenic controls (Fig. S14B-D). Together, these lines of evidence suggest that PRC2.1 mediates the effects of a high sugar diet on sweet taste by directly controlling the expression of transcription factors that regulate a transcriptional program required for sweet taste.

The persistent phenotypic memory of the food environment is dependent on PRC2.1

The cellular fates created by Polycomb Group proteins are inherited as stable memories across cell divisions to ensure phenotypic stability even in the absence of the triggering stimuli (Laprell et al., 2017) (Coleman and Struhl, 2017) (Audergon et al., 2015) (Ragunathan et al., 2015) (Gaydos et al., 2014). We therefore asked if the neural state created by PRC2.1 in the high sugar diet environment was maintained when flies were moved to the control diet after eating a sugar diet for 7 days (SD>CD) (Fig. 6A) We found that these animals had lower sweet taste, similar to that of age-matched flies fed a sugar diet for 7 days (Fig.6B, SD>CD compared to CD>SD). However, their triglyceride levels were similar to those of control diet flies (Fig. S14E), suggesting that while fat storage was reversible when flies only had access to the “healthy” control diet, sweet taste deficits persisted.

To understand how this phenotype compares to that of the control diet flies at the molecular level, we conducted TRAP of the Gr5a+ neurons of flies in the SD>CD and CD>CD conditions. mRNAs from flies on a SD>CD showed a strong negative skew in log₂ fold changes compared to the control diet group (Fig. 6C, -2.02), reminiscent of the skew we observed in flies fed a sugar diet (Fig. 4C). Furthermore, we observed that 47% (310/658) of genes in the
transcriptional network repressed by PRC2.1 on a sugar diet were still decreased in SD>CD flies (Fig. 6D). Interestingly, the SD>CD animals clustered with the sugar diet 7 (SD7) group compared to sugar diet 3 (SD3) and Pcl mutants fed a sugar diet (Fig. 6D). Thus, at the molecular level, half of the neural state established by dietary sugar via PRC2.1 persisted. In addition to the GO term categories changed in the sugar diet condition (like signal transduction, cilium assembly, detection of chemical stimulus, carbohydrate metabolic process), we also found new GO term categories present (Fig. S15), suggesting that there may be aspects of this persistent state that are also novel. To test the hypothesis that PRC2.1 plays an active role in maintaining this neural state, we inhibited PRC2 activity during the “recovery” diet using the EEDi inhibitor (SD>CD+EEDi). Remarkably, these animals showed a restoration of wild-type sweet taste (Fig. 6E, green triangles). Together, these data indicate that the sensory neurons retain a phenotypic memory of the sugar diet environment, and that PRC2.1 is constitutively required for its persistence.

Discussion

In this study we set out to understand how the food environment alters the gustatory system as a model to define how interactions between genes and environment persistently reprogram complex behavior. Specifically, we took advantage of the ability of diet to change taste sensation to identify the molecular mechanisms through which the food environment changes neural state, physiology, and behavior in a circuit-specific way. Here we show that the decrease in sweet taste sensation that flies experience after chronic exposure to a high sugar diet is caused by the cell-autonomous action of the Polycomb Repressive Complex 2.1 in the sweet gustatory neurons. Mutations and pharmacological inhibition of PRC2.1 blocked the effects of the food environment on neural activity, behavior, and obesity. While we do not
exclude the possibility that PRC1 and PhoRC may also be involved, we found that mutations or knockdown in these complexes had no effect on taste. In the high sugar food environment, PRC2.1 occupancy was redistributed, leading to the repression of transcription factors, neural, signaling, and metabolic genes that decreased the responsiveness of the Gr5a+ neurons and the fly’s sensory experience. However, we discovered that PRC2.1 did not directly bind to neuronal genes in these cells and that, instead, it targeted transcription factors involved in sensory neuron development, synaptic function, and axon targeting. Specifically, on a high sugar diet Pcl binding was increased at the cad, GATAe, nub/pdm, Ptx1 loci and decreased at the Scro locus, with corresponding changes in the mRNA levels of these genes (Fig. 6E, model). Together, the decrease in the mRNA levels of the 4 activators (cad, GATAe, nub/pdm, Ptx1) and the increase in the repressor (Scro), led to the repression of 658 genes implicated in synaptic function, signal transduction, and metabolism. Interestingly, we uncovered several feed-forward regulatory loops among the transcriptional regulators, suggesting that this hub could ensure the stability of the neural state unless the change in environment is chronic. This transcriptional network controlled by these regulators was necessary for sweet taste, because knockdown of the four activators and a few of their targets, as well as overexpression of Scro (Fig. 5B and Fig. S14B-D), resulted in a decrease in sweet taste on a control diet.

Interestingly, several of the transcription factors we identified – Ptx1, Scro, and nub/pdm – have been shown to control the proper branching, synaptic connectivity, and function of sensory neurons (Corty et al., 2016; Iyer et al., 2013; Neumann and Cohen, 1998; Parrish et al., 2006, 2007; Vorbrüggen et al., 1997; Zaffran et al., 2000), while others (cad, nub/pdm) play a role in neuroblast development (Doe, 2017; Kohwi and Doe, 2013); PRC2 also functions as a competence factor in neural proliferation, differentiation and sensory neurons (Bahrampour et al., 2019; Doe, 2017; Parrish et al., 2007). Importantly, we found that the 4 activators that are
repressed by Pcl in the high sugar condition are enriched in the Gr5a+ cells on a control diet, while Scro is depleted. Thus, the transcriptional network of ~658 genes controlled by these transcription factors may define the intrinsic properties of the sweet sensing neurons. Interestingly, in addition to there being substantial overlap in the gene batteries regulated by the 5 transcription factors, we also found that these target genes were functionally interconnected in a network, especially those involved in signaling, synaptic function, and cell adhesion such as the kinase haspin, the adenylate cyclase ACXD, sytalpha, Arc1, tetraspanin, jonan, and innexin proteins among others, which may affect the circuit both at the functional and connectivity levels. Since we did not observe a change in the expression of the sweet taste receptors, or the misexpression of other taste receptors (Scott, 2018), our data are not consistent with a complete loss of identity of the Gr5a+ neurons. Instead, we propose that PRC2.1 tunes the sweet sensory neurons to the environment by altering a transcriptional network that controls the intrinsic properties of these cells, especially those involved in signal transduction, connectivity, synaptic function, and metabolism. Studies that test the effects of this network on the connectivity, morphology, and signal transduction of the sweet sensory neurons will shed light on how exactly the transcriptional remodeling caused by PRC2.1 impacts the Gr5a+ cells.

A previous study showed that PRC2 was required to maintain the properties of medium spiny neurons in post-mitotic cells (von Schimmelmann et al., 2016), but here we found that it was its active redistribution in response to the environment – rather than its loss – that altered the properties of these cells. Thus, our work opens up the exciting possibility that PRC2 may modulate neural plasticity in response to environmental conditions. In other post-mitotic cells, PcG proteins alter transcriptional programs according to environmental stressors, such as oxidative stress, injury response, temperature, and light (Kolybaba and Classen, 2014; Marasca
et al., 2018). Our findings are in line with these and contend that, in addition to establishing cell fates, PRC2 regulates “neural states” depending on the environment.

Using neuroepigenetic mechanisms to tune neural states to external conditions could provide several advantages compared to the medley of other cellular, receptor, or synaptic plasticity based mechanisms. Specifically, it would allow cells to 1) orchestrate a coordinated response, 2) create a memory of the environment, and 3) buffer small fluctuations until a substantial challenge is perceived. It is particularly fascinating to think about the molecular mechanisms through which these neural states may be established. The need of neurons to constantly maintain their identity may mean that environmental signals like the extent of sensory stimulation could alter the expression of developmental gene batteries and affect neural states (Deneris and Hobert, 2014). Indeed, it has been speculated that some forms of plasticity may re-engage developmental programs that specify the intrinsic properties of neurons (Marder and Prinz, 2002; Parrish et al., 2014). As mentioned above, we observed that the regulators of the transcriptional network we uncovered function in sensory neuron development and are enriched in the Gr5a+ cells. Thus, it could be a hallmark of neuroepigenetic plasticity to exploit developmental programs, linking the known role of PRC2 in establishing cell fates with this newly discovered function in modulating cell states. Incidentally, engaging developmental programs could be the reason why some environments and experiences leave a memory that leads to the persistent expression of the phenotype beyond the presence of the triggering stimulus, as these could target neural connectivity and set synaptic weights thresholds. Defining the circuit-specific changes of the pioneering studies on maternal care, addiction, and learning and memory (Champagne, 2008; Dulac, 2010; Nestler and Lüscher, 2019; Sweatt, 2013) would put this hypothesis to test. Here we found that the changes in taste sensation and half of the sugar diet neural state set by PRC2.1 remained even after animals were moved back to the
control diet for a week. A limitation of our study is that due to the small number of Gr5a+ neurons and their anatomically inaccessible location, we were not able to measure the identity of the molecular memory in these cells alone. However, we saw that the phenotypic memory of the high sugar food environment was dependent on the constitutive action of PRC2.1. Based on other studies showing that the H3K27 methyl mark acts as a molecular memory during development (Coleman and Struhl, 2017; Laprell et al., 2017), we speculate that this is likely to be the molecular signal in the Gr5a+ cells too. It is interesting to note that a long term (4 weeks) increase in H3K27 methylation at the BDNF exon III and IV promoter was measured in response to social defeat, although the role of this mark or PRC2 in the behavioral phenotype or its persistence was not examined (Tsankova et al., 2006). The question of how exactly PRC2.1 senses the changes in the food environment also remains open. Recent studies suggest that the activity of Polycomb Group proteins is directly and indirectly linked to cellular metabolism, including kinases signaling cascades, GlcNAcylation, and the availability of cofactors for histone modifications, (Kolybaba and Classen, 2014; Marasca et al., 2018). Our experiments show that PRC2.1 binding changes depending on the dietary environment, and provides a new model to tackle this question.

In conclusion, we show that PRC2.1 mediates the effects of dietary sugar on sweet taste by establishing persistent alterations in the taste neurons that remain as a phenotypic and neural memory of the previous food environment. Given the importance of taste in modulating food intake, we speculate that this memory may lock animals into patterns of feeding behavior that become maladaptive and promote obesity, and, indeed, we found that Pcl mutant flies are protected from diet-induced obesity. Thus, the food environment, like other experiences, can induce lasting alterations that restrict the behavioral plasticity of animals and impact disease risk. Since the content of sugar in processed foods is similar or higher than that we fed flies and
the function of PcG proteins is conserved from plants to humans, our work is broadly relevant to understanding the effects of the food environment on a whole range of diet-related conditions and diseases that affect billions of people worldwide.
Methods

Fly Husbandry and Strains:
All flies were grown and maintained on cornmeal food (Bloomington Food B recipe) at 25°C and 45%–55% humidity under a 12:12 hour light-dark cycle (ZT0 at 9 AM). Male flies were collected under CO2 anesthesia 1-3 days after eclosion and maintained in a vial that housed 35-40 flies. Flies were acclimated to their new vial environment for an additional 2 days. For all experiments, flies were changed to fresh food vials every other day.

For all dietary manipulations, the following compounds were mixed into standard cornmeal food (Bloomington Food B recipe) (0.58 calories per gram) by melting, mixing, and pouring new vials as in (Musselman and Kühnlein, 2018) and (Na et al., 2013). For the 30% sugar diet (1.41 calories per gram) Domino granulated sugar w/v was added. For the EED226 inhibitor diet (AxonMedchem), EED226 was solubilized in 10% DMSO and added to control or 30% sugar diet at a total concentration of 8 uM.

For genetic manipulations the GAL4/UAS system was used to express transgenes of interest in *Gustatory receptor 5a* Gr5a-GAL4. For each GAL4/UAS cross, transgenic controls were made by crossing the w1118CS (gift from A. Simon) to GAL4 or UAS flies, sex-matched to those used in the GAL4/UAS cross.

The following fly lines are used in this paper: Gr5a-GAL4/Cyo (gift from K. Scott, University of California, Berkeley), nsyb-GAL4/Cyo (gift from J. Simpson, University of California, Santa Barbara), Gr64f-GAL4/Cyo (gift from H. Amrein, University of Texas, A&M), UAS-GCaMP6s-Brp-mCherry (BDSC #77131), Tubulin-GAL80s (BDSC #7018), UAS-Ogt-RNAi and UAS-Ogt-GFP/Cyo (gift from C. Lehner, University of Zurich), UAS-Pcl (FlyORF #F001897), UAS-Pcl-RNAi (VDRC #v220046, BDSC #33945), UAS-Rpl3-3XFLAG (gift from D. Dickman, University of Southern California), Pcl+/+/CKG (gift from G. Cavalli, Université de Montpellier),
Pcl\textsuperscript{241}/Cyo, E(z)\textsuperscript{1249}/TM3, Su(z)\textsuperscript{12} \textsuperscript{253}, Esc\textsuperscript{289} (gift from N.Liu, Chinese Academy of Sciences), Pc\textsuperscript{1}/TM1 (BDSC #1728), ph-d\textsuperscript{601} ph-p\textsuperscript{602}/FM7 (BDSC #5444), Psc\textsuperscript{h27}/Cyo (BDSC #5547), UAS-LT3-Dam (gift from A.H Brand, University of Cambridge), UAS-LT3-Dam::PCL (made in this paper), UAS-Arc1-RNAi (BDSC #25954), Arc\textsuperscript{esm13} (BDSC #37530), Arc1\textsuperscript{esm18} (BDSC #37531), UAS-Arc1 (BDSC #37532), UAS-Ptx1-RNAi (VDRC #107785), UAS-cad-RNAi (BDSC #34702), UAS-GATAe-RNAi (BDSC #33748), UAS-nub (FlyORF #F000147), UAS-Scro-RNAi (BDSC #29387), UAS-Ptx1 (FlyORF #F003469), UAS-Scro (FlyORF F003427), UAS-nub-RNAi (BDSC #55305), UAS-cad (FlyORF #F000471) UAS-Jarid2-RNAi (BDSC #32891, #26184), UAS-ACXD-RNAi (BDSC #62871, #35589), UAS-jing-RNAi (BDSC #27084), UAS-Sfmbt-RNAi (BDSC #32473).

**Proboscis Extension Response:**

Male flies were food deprived for 18-24 hours in a vial with a Kimwipe dampened with 2 mL of milliQ-filtered deionized (milliQ DI) water. Proboscis extension response (PER) was carried out as described in (Shiraiwa and Carlson, 2007).

**Proboscis Immunofluorescence:**

Probosces were dissected in 1xPBS and fixed in 4% PFA, mounted in FocusClear (CelExplorer) on coverslips. Cell bodies were imaged using a FV1200 Olympus confocal with a 20x objective. Cells were counted using Imaris Image analysis software.

**Triglyceride Measurements:**

Total triglycerides normalized to total protein were measured as described in (Tennesen et al., 2014). Briefly, two flies per biological replicate were homogenized in lysis buffer (140 mM NaCl,
50 mM Tris-HCl pH 7.4, 0.1% Triton-X) containing protease inhibitor (Thermo Scientific). Lysate was used to determine protein and triglyceride concentrations using Pierce BCA assay (Thermo Scientific, abs 562 nm) and Triglycerides LiquiColor test (Stanbio, abs 500 nm), respectively.

**Calcium Imaging:**
Male flies expressing GCaMP6s-Brp-mCherry (Kiragasi et al., 2017) in the sweet sensing neurons were food deprived for 18-24 hours. The flies for live imaging were prepared similar to LeDue et al. (LeDue et al., 2015). Briefly, flies were fixed to a custom-printed plastic slide with paraffin wax and the proboscis waxed to an extended position. Distal leg segments were removed to prevent tarsal interference with labellar stimulation. To image the SEZ, sugarless artificial hemolymph solution filled the well surrounding the head. Subsequently, the dorsal cuticle between the eyes was removed by microdissection to expose the brain. Each fly proboscis was tested with milliQ water before stimulating with 20% sucrose dissolved in milliQ water. Stimulus (a piece of Kimwipe soaked in tastant and held with forceps) delivery to the proboscis was manual and timed to coincide with the 100th recording sample of each time series. Imaging was carried out using an upright confocal microscope (Olympus, FluoView 1200 BX61WI), a 20x water-immersion objective and laser excitation at 488 and 543 nm. Recordings were made at 4 Hz (512 x 512 pixels). Plane of interest was kept to the most ventral neuropil regions innervated by the sweet sensing neurons.

**RNA Extraction and Quantitative RT-PCR:**
For all RNA extractions used for qPCR, fly heads from 10-20 flies were dissected into Trizol (Ambion) and homogenized with plastic pestles. RNA was extracted by phenol chloroform (Ambion), and precipitated by isopropanol with Glycobule Coprecipitant (Invitrogen). RNA pellet
was washed as needed with 75% ethanol and subsequently eluted in nuclease free water and treated with DNase I, according to manufacturer's instructions (Turbo DNA-free DNA removal kit, Ambion). All steps were carried out in RNase free conditions, and RNA was stored at -80°C until further processing.

Complementary DNA was synthesized by Superscript III (Invitrogen) reverse transcriptase with the addition of Ribolock RNase inhibitor (Thermo Fisher Scientific). qPCR reactions were carried out using Power SYBR Green PCR master mix (Applied Biosystems) based on manufacturer's instructions. Primers were added at a 2.5uM concentration. All reactions were run on a 96-well plate on the StepOnePlus Real-Time PCR System (Applied Biosystems) and quantifications were made relative to the reference gene *Ribosomal protein 49 (Rp49).* Primer sequences are listed below and were tested for efficiency prior to the qPCR experiment:

| Gene                  | Sequence                                      |
|-----------------------|-----------------------------------------------|
| Rp49 Forward          | ATGCTAAGCTGTCGCACAAA                           |
| Rp49 Reverse          | ACTTCTGAATCCGTTGGGC                            |
| Pcl Forward           | ACCCATGAAAATGCAACAGCC                          |
| Pcl Reverse           | TTATCCGCATGGTGAGGCAAA                         |
| Pcl_Gibson_Foward     | tctctgaagagatctgGCCGGCGCaATGATGAA CAACCATTTCACCTGCAAC |
| Pcl_Gibson_Reverse    | agtaagttcctcacaagatctcTCAATTTCACAAG CAATCCAATCGCC |

**Affinity purification of ribosome associated mRNA (TRAP):**

300 heads (10,000 cells) per biological replicate were collected using pre-chilled sieves in liquid nitrogen on dry ice. Frozen heads were then lysed as previously described (Chen and Dickman,
From the lysate total RNA was extracted by TRIzol LS Reagent (ThermoFisher scientific, 10296010) as input. The remainder of the lysate was incubated with Dynabeads protein G (ThermoFisher scientific, 10004D) to preclear samples for 2 hours and subsequently incubated with Dynabeads protein G coated with an anti-Flag antibody (Sigma, F1804). The lysate-beads mixture was incubated at 4°C with rotation for 2 hours, then washed with buffer. RNA was extracted from ribosomes bound to the beads by TRIzol Reagent (Chen and Dickman, 2017).

**Targeted DNA Adenine Methyltransferase Identification (TaDa):**

For the UAS-LT3-Dam::Pcl construct, the coding region of the Pcl gene was amplified from the pCRE-NDAM-Myc-DO69-Pcl (gift from Bas Van Steensel, The Netherlands Cancer Institute) with primers listed above and assembled into the UAS-LT3-DAM plasmid (gift from Andrea Brand, University of Cambridge) using the NEBuilder® HiFi DNA Assembly kit based on manufacturers instructions (NEB). Transgenic animals were validated by RT-PCR. TaDa lines were crossed to Gr5a-GAL4; tubulin-GAL80ts. All animals were raised and maintained at 20 °C. Dam::Pcl and Dam was induced at 28 °C for 18-20 hours. For all experiments 300 heads of males and female flies were collected per replicate on dry ice by sieving. DNA was extracted from frozen heads following kit instructions (Invitrogen). For identification of methylated regions purified DNA was digested by DpnI followed by PCR purification of digested sequences. TaDa adaptors were ligated by T4 DNA ligase (NEB). Adapter ligated DNA was PCR amplified according to protocol (Marshall et al., 2016), and subsequently purified. Purified DNA was digested with DpnII followed by sonication to yield fragments averaging 200-300bp. TaDa adaptors were removed from sonicated DNA by digestion (Marshall et al., 2016).

**Library Preparation for TRAP and TaDa:**
For RNA sequencing libraries were generated using the Ovation SoLo RNA-Seq System for *Drosophila* (Nugen, 0502-96). All reactions included integrated HL-dsDNase treatment (ArcticZymes, Cat. #70800-201). For DNA sequencing libraries were generated using the Takara ThruPlex kit (cat #022818) using 3ng input and 10 cycles of PCR. All libraries were sequenced on the Illumina NextSeq platform (High-output kit v2 75 cycles) at the University of Michigan core facility.

**High Throughput RNA-seq Analysis**

Fastq files were assessed for quality using FastQC (Andrews and Others, 2010). Reads with a quality score below 30 were discarded. Sequencing reads were aligned by STAR (Dobin et al., 2013) to *dmel*-all-chromosomes of the dm6 genome downloaded from Ensemble, and gene counts were obtained by HTseq (Anders et al., 2015). Count files were used as input to call differential RNA abundance by DESeq2 (Love et al., 2014). All pairwise comparisons were made to the control diet of the corresponding genotypes, such that sugar diet three days and sugar diet seven days were compared to the age matched control diet group. In *Pcl* mutants experiments, the pairwise comparison was made between sugar diet and control diet within the age-matched *Pcl*429 genotype group. A cutoff of qval<0.1 was used to call differentially expressed genes. Skew in log2 fold changes was measured using the R package Skewness (e1071). RNAseq data visualization was carried out in R studio using ggplot2 and the following packages, pheatmap (Kolde, 2012), Venn euler (Wilkinson, 2012), and EnhancedVolcano (Blighe, 2019). To cluster columns and rows in pheatmap "Ward.D " clustering was used.

**High Throughput TaDa and CATaDa Analysis**
Fastq files were assessed for quality using FastQC (Andrews and Others, 2010). Reads with a quality score below 30 were discarded. The damidseq_pipeline was used to align, extend, and generate log2 ratio files \((\text{Dam::Pcl/Dam})\) in GATC resolution as described previously (Marshall and Brand, 2015). In short, the pipeline uses Bowtie2 (Langmead and Salzberg, 2012) to align reads to \(dme\)-all-chromosomes of the dm6 genome downloaded from Ensemble, followed by read extension to 300 bp (or to the closest GATC, whichever is first). Bam output is used to generate the ratio file in bedgraph format. Bedgraph files were converted to bigwig and visualized in the UCSD genome browser. Correlation coefficients and PCA plot between biological replicates were computed by multibigwigSummary and plotCorrelation in deepTools (Ramírez et al., 2016). Fold Change traces for SD/CD of \(\log_2(\text{Dam::Pcl/Dam})\) were generated by calculating the mean profile of all replicates for each condition and subsequently calculating fold change between the sugar diet and control diet condition with deepTools bigwigCompare (Ramírez et al., 2016). Peaks were identified from \(\log_2(\text{Dam::Pcl/Dam})\) ratio files using find_peaks (FDR<0.01) (Marshall and Brand, 2015). To do this, the binding intensity thresholds are identified from the dataset, the dataset is then shuffled randomly, and the frequency of consecutive regions (i.e. GATC fragments or bins) with a score greater than the threshold is calculated. The FDR is the observed / expected for a number of consecutive fragments above a given threshold. Association of genes to peaks was made using the peaks2genes script (Marshall and Brand, 2015) and dm6 genome annotations. Overlapping intervals or nearby intervals were merged into a single interval using MergeBED in Bedtools (Quinlan and Hall, 2010). Intervals common in all replicate peak files were identified by Multiple Intersect in Bedtools (Quinlan and Hall, 2010). DiffBind was used to determine differentially bound sites on peak files in bed format based on differences in read intensities (Stark et al., 2011).
For CATaDa experiments, all analyses were performed similar to those of TaDa with the exceptions that 1) Dam only profiles were not normalized as ratios but shown as non normalized binding profiles, 2) Dam only coverage plots were generated by converting bam files to bigwig files normalized to 1x dm6 genome, and 3) peaks were called using MACS2 call peaks on alignment files without building the shifting model with an of FDR<0.05 (Feng et al., 2012). To determine the proportion of genes that fit within the various chromatin domain subtypes, we first matched Dam::Pcl/Dam targets to coordinates identified by Filion et al and then determined their gene count in each chromatin subtype (observed) compared to the whole genome (expected).

**iPAGE Analysis**

All Gene Ontology (GO) term enrichment analysis was performed using the iPAGE package (Goodarzi et al., 2009), using gene-GO term associations extracted from the Flybase dmel 6.08 2015_05 release (Thurmond et al., 2019). For analysis of ribotrap data, iPAGE was run in continuous mode, with log₂ fold changes divided into seven equally populated bins. For analysis of DamID and TF regulatory targets, iPAGE was run in discrete mode, using the groups specified for each calculation. For all discrete calculations, independence filtering was deactivated due to the less informative available signal. All other iPAGE settings were left at their default values. All shown GO terms pass the significance tests for overall information described in (Goodarzi et al., 2009); in addition, for each term, individual bins showing especially strong contributions (p-value such that a Benjamini-Hochberg FDR (Benjamini and Hochberg, 1995) calculated across that row, yields q<0.05) are highlighted with a strong black box.
**PREdictor**

Identification of predicted PRE sites was performed exactly as described in (Khabiri and Freddolino, 2019), using the dm6 genome. As suggested in (Khabiri and Freddolino, 2019), we use a threshold confidence score of 0.8 to identify the PREs used in the present analysis. A complete list of predicted PREs, with accompanying confidence scores, is shown in supplementary file 1. Enrichments of overlap between different PRE classes and Pcl occupancy locations were calculated by comparing the observed overlap frequency with the overlaps for 1,000 random shufflings of the binding/differential binding peak locations (calculated using bedtools 2.17.0 (Quinlan and Hall, 2010).

**Calculation of regulatory targets of transcription factors**

To identify likely targets of each transcription factor of interest, we drew upon the transcription factor binding site calculations described in (Khabiri and Freddolino, 2019), in which the motif of each transcription factor was scanned along every base pair of the *D. melanogaster* dm6 genome using FIMO (Grant et al., 2011), and the base pair-wise binding results converted to robust z-scores. For each TF, we then considered its regulon to consist of all genes with at least one binding site with z-score above a TF-specific threshold within 2 kb upstream of the beginning of the gene. We identified TF-specific thresholds by manual inspection of a plot of the average expression changes between conditions vs. threshold, aiming to identify a point of maximum information content relative to noise (see supplemental figure 8 for the plots used to identify TF-specific z-score thresholds). Once the set of targets (regulon) for each factor was identified, we tested for significant enrichment or depletion of overlaps between the regulons using Fisher’s exact test, reporting Benjamini-Hochberg false discovery rates (FDRs) (Benjamini
and Hochberg, 1995). All calculated odds ratios were positive, indicating enriched overlaps between the regulons.

**STRING network analysis**

To develop a functional network between the candidate neural targets of *cad, Ptx1, nub, GATAe* and *Scro*, we uploaded genes categorized into neural/signaling GO terms based on DAVID functional annotations (Huang et al., 2008) (Huang et al., 2009) to the STRING database search (Szklarczyk et al., 2019). Genes were clustered by their reported protein-protein interactions and corresponding confidence scores (Szklarczyk et al., 2019) and plotted in Cytoscape (v3.7.1) (Shannon et al., 2003). In this network edges do not represent direct protein protein interaction but rather represent a functional interaction. For network see Supplementary file 5.

**Data Analysis and Statistics**

Statistical tests, sample size, and p or q values are listed in each figure legend. Data were evaluated for normality and appropriate statistical tests applied if data were not normally distributed, all the tests, biological samples, and the p and q values are listed in the figure legends and specific analysis under each methods session. Because the inferential value of a failure to reject the null hypothesis in frequentist statistical approaches is limited, for all RNA-seq expression datasets, we coupled our standard differential expression with a test for whether each gene could be flagged as ‘significantly not different’. Defining a region of practical equivalence (ROPE) as a change of no more than 1.5-fold in either direction, we tested the null hypothesis of at least a 1.5-fold change for each gene, using the gene-wise estimates of the standard error in log2 fold change (reported by Deseq2) and the assumption that the actual log2 fold changes are normally distributed. Rejection of the null hypothesis in this test is taken as
positive evidence that the gene’s expression is not changed substantially between the conditions of interest. Python code for the practical equivalence test can be found on Github as calc_sig_unchanged.py. All data in the figures are shown as Mean ± SEM, **** p < 0.0001, *** p < 0.001, ** p < 0.01,*p< 0.05 unless otherwise indicated.

Data availability

All high throughput sequencing data files can be found on Gene Expression Omnibus GSE146245.

Conflict of interests

The authors declare no conflict of interest.

Acknowledgements

We thank Sundeep Kalantry, Josie Clowney, and Giacomo Cavalli for helpful comments and discussions. We also thank the University of Indiana at Bloomington, the VDRC, the FLYORF stock collections, and all the investigators who kindly shared fly lines with us. We also thank Daniel Wilinski for help with data analysis, Dion Dickman and Xun Chen for helpful comments on riboTRAP, and Margarita Brovkina (Clowney lab) for helpful discussions on TaDa analysis. Matt Burkhard assisted with analysis of expression changes for different Gene Ontology terms. Julia Kuhn designed some of the graphics for the manuscript. This work was funded by NIH R00 DK-97141 and NIH 1DP2DK-113750, the Klingenstein-Simons Fellowship in the Neurosciences, and the Rita Allen Foundation (to M.D.) and NIH R35 GM-128637 to (P.L.F).

Author Contributions
A.V performed all experiments and analysed all RNAseq, TaDa, and CATaDa datasets, with the exception of in vivo calcium imaging. B.T.G helped with PER and triglyceride measurements. C.E.M performed in vivo calcium imaging. M.K and P.L.F developed the PREdictor, calculated regulatory targets of transcription factors, ran and analyzed Gene Ontology term enrichment analysis, provided statistical consultation, and tested for practical significance. M.D oversaw the project and secured funding. A.V and M.D designed the experiments, wrote the manuscript, and prepared the figures with input from all authors.
References

Ahart, Z.C., Martin, L.E., Kemp, B.R., Banik, D.D., Roberts, S.G.E., Torregrossa, A.M., and Medler, K. (2019). Differential effects of diet and weight on taste responses in diet-induced obese mice.

Allis, C.D., and Jenuwein, T. (2016). The molecular hallmarks of epigenetic control. Nat. Rev. Genet. 17, 487–500.

Anders, S., Pyl, P.T., and Huber, W. (2015). HTSeq--a Python framework to work with high-throughput sequencing data. Bioinformatics 31, 166–169.

Andrews, S., and Others (2010). FastQC: a quality control tool for high throughput sequence data.

Ashley, J., Cordy, B., Lucia, D., Fradkin, L.G., Budnik, V., and Thomson, T. (2018). Retrovirus-like Gag Protein Arc1 Binds RNA and Traffics across Synaptic Boutons. Cell 172, 262–274.e11.

Audergon, P.N.C.B., Catania, S., Kagansky, A., Tong, P., Shukla, M., Pidoux, A.L., and Allshire, R.C. (2015). Epigenetics. Restricted epigenetic inheritance of H3K9 methylation. Science 348, 132–135.

Aughey, G.N., Gomez, A.E., Thomson, J., Yin, H., and Southhall, T.D. (2018). CATaDa reveals global remodelling of chromatin accessibility during stem cell differentiation in vivo. eLife, 7.

Bahrampour, S., Jonsson, C., and Thor, S. (2019). Brain expansion promoted by polycomb-mediated anterior enhancement of a neural stem cell proliferation program. PLoS Biol. 17, e3000163.

Beauchamp, G.K., and Jiang, P. (2015). Comparative biology of taste: Insights into mechanism and function. Flavour 4, 9.

Benjamini, Y., and Hochberg, Y. (1995). Controlling the False Discovery Rate: A Practical and Powerful Approach to Multiple Testing. J. R. Stat. Soc. Series B Stat. Methodol. 57, 289–300.

Bertino, M., Beauchamp, G.K., and Engelman, K. (1982). Long-term reduction in dietary sodium alters the taste of salt. Am. J. Clin. Nutr. 36, 1134–1144.

Bligh, K. (2019). EnhancedVolcano: Publication-ready volcano plots with enhanced colouring and labeling. R package version 1.2. 0.

Campbell, R.R., and Wood, M.A. (2019). How the epigenome integrates information and reshapes the synapse. Nat. Rev. Neurosci. 20, 133–147.

Champagne, F.A. (2008). Epigenetic mechanisms and the transgenerational effects of maternal care. Front. Neuroendocrinol. 29, 386–397.
Chan, C.S., Rastelli, L., and Pirrotta, V. (1994). A Polycomb response element in the Ubx gene that determines an epigenetically inherited state of repression. EMBO J. 13, 2553–2564.

Chen, X., and Dickman, D. (2017). Development of a tissue-specific ribosome profiling approach in Drosophila enables genome-wide evaluation of translational adaptations. PLoS Genet. 13, e1007117.

Chen, K., Yan, J., Suo, Y., Li, J., Wang, Q., and Lv, B. (2010). Nutritional status alters saccharin intake and sweet receptor mRNA expression in rat taste buds. Brain Res. 1325, 53–62.

Chyb, S., Dahanukar, A., Wickens, A., and Carlson, J.R. (2003). Drosophila Gr5a encodes a taste receptor tuned to trehalose. Proc. Natl. Acad. Sci. U. S. A. 100 Suppl 2, 14526–14530.

Coleman, R.T., and Struhl, G. (2017). Causal role for inheritance of H3K27me3 in maintaining the OFF state of a Drosophila HOX gene. Science 356.

Corty, M.M., Tam, J., and Grueber, W.B. (2016). Dendritic diversification through transcription factor-mediated suppression of alternative morphologies. Development 143, 1351–1362.

Crow, J.M. (2012). Obesity: insensitive issue. Nature 486, S12–S13.

Deneris, E.S., and Hobert, O. (2014). Maintenance of postmitotic neuronal cell identity. Nat. Neurosci. 17, 899–907.

Dobin, A., Davis, C.A., Schlesinger, F., Drenkow, J., Zaleski, C., Jha, S., Batut, P., Chaisson, M., and Gingeras, T.R. (2013). STAR: ultrafast universal RNA-seq aligner. Bioinformatics 29, 15–21.

Doe, C.Q. (2017). Temporal Patternining in the Drosophila CNS. Annu. Rev. Cell Dev. Biol. 33, 219–240.

Dulac, C. (2010). Brain function and chromatin plasticity. Nature 465, 728–735.

Feng, J., Liu, T., Qin, B., Zhang, Y., and Liu, X.S. (2012). Identifying ChIP-seq enrichment using MACS. Nat. Protoc. 7, 1728–1740.

Filion, G.J., van Bemmel, J.G., Braunschweig, U., Talhout, W., Kind, J., Ward, L.D., Brugman, W., de Castro, I.J., Kerkhoven, R.M., Bussemaker, H.J., et al. (2010). Systematic protein location mapping reveals five principal chromatin types in Drosophila cells. Cell 143, 212–224.

Gaydos, L.J., Wang, W., and Strome, S. (2014). Gene repression. H3K27me and PRC2 transmit a memory of repression across generations and during development. Science 345, 1515–1518.

Glendinning, J.I., Domdom, S., and Long, E. (2001). Selective adaptation to noxious foods by a herbivorous insect. J. Exp. Biol. 204, 3355–3367.

Goodarzi, H., Elemento, O., and Tavazoie, S. (2009). Revealing global regulatory perturbations
across human cancers. Mol. Cell 36, 900–911.

Grant, C.E., Bailey, T.L., and Noble, W.S. (2011). FIMO: scanning for occurrences of a given motif. Bioinformatics 27, 1017–1018.

Huang, D.W., Sherman, B.T., and Lempicki, R.A. (2008). Systematic and integrative analysis of large gene lists using DAVID Bioinformatics Resources. Nature Protoc. 2009; 4 (1): 44--57.

Huang, D.W., Sherman, B.T., and Lempicki, R.A. (2009). Bioinformatics enrichment tools: paths toward the comprehensive functional analysis of large gene lists. Nucleic Acids Res. 37, 1–13.

Iyer, E.P.R., Iyer, S.C., Sullivan, L., Wang, D., Meduri, R., Graybeal, L.L., and Cox, D.N. (2013). Functional Genomic Analyses of Two Morphologically Distinct Classes of Drosophila Sensory Neurons: Post-Mitotic Roles of Transcription Factors in Dendritic Patterning. PLoS ONE 8, e72434.

Kaufman, A., Choo, E., Koh, A., and Dando, R. (2018). Inflammation arising from obesity reduces taste bud abundance and inhibits renewal. PLoS Biol. 16, e2001959.

Khabiri, M., and Freddolino, P.L. (2019). Genome-wide Prediction of Potential Polycomb Response Elements and their Functions.

Kiragasi, B., Wondolowski, J., Li, Y., and Dickman, D.K. (2017). A Presynaptic Glutamate Receptor Subunit Confers Robustness to Neurotransmission and Homeostatic Potentiation. Cell Rep. 19, 2694–2706.

Kohwi, M., and Doe, C.Q. (2013). Temporal fate specification and neural progenitor competence during development. Nat. Rev. Neurosci. 14, 823–838.

Kolde, R. (2012). Pheatmap: pretty heatmaps. R Package Version.

Kolybaba, A., and Classen, A.-K. (2014). Sensing cellular states—signaling to chromatin pathways targeting Polycomb and Trithorax group function. Cell Tissue Res. 356, 477–493.

Langmead, B., and Salzberg, S.L. (2012). Fast gapped-read alignment with Bowtie 2. Nat. Methods 9, 357–359.

Laprell, F., Finkl, K., and Müller, J. (2017). Propagation of Polycomb-repressed chromatin requires sequence-specific recruitment to DNA. Science 356, 85–88.

Laugesen, A., Højfeldt, J.W., and Helin, K. (2019). Molecular Mechanisms Directing PRC2 Recruitment and H3K27 Methylation. Mol. Cell 74, 8–18.

LeDue, E.E., Chen, Y.-C., Jung, A.Y., Dahanukar, A., and Gordon, M.D. (2015). Pharyngeal sense organs drive robust sugar consumption in Drosophila. Nature Communications 6.

Love, M.I., Huber, W., and Anders, S. (2014). Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol. 15, 550.
Maliphol, A.B., Garth, D.J., and Medler, K.F. (2013). Diet-induced obesity reduces the responsiveness of the peripheral taste receptor cells. PLoS One 8, e79403.

Marasca, F., Bodega, B., and Orlando, V. (2018). How Polycomb-Mediated Cell Memory Deals With a Changing Environment: Variations in PcG complexes and proteins assortment convey plasticity to epigenetic regulation as a response to environment. Bioessays 40, e1700137.

Marder, E., and Prinz, A.A. (2002). Modeling stability in neuron and network function: the role of activity in homeostasis. Bioessays 24, 1145–1154.

Marshall, O.J., and Brand, A.H. (2015). damidseq_pipeline: an automated pipeline for processing DamiD sequencing datasets. Bioinformatics 31, 3371–3373.

Marshall, O.J., Southall, T.D., Cheetham, S.W., and Brand, A.H. (2016). Cell-type-specific profiling of protein–DNA interactions without cell isolation using targeted DamiD with next-generation sequencing. Nat. Protoc. 11, 1586–1598.

May, C.E., Vaziri, A., Lin, Y.Q., Grushko, O., Khabiri, M., Wang, Q.-P., Holme, K.J., Pletcher, S.D., Freddolino, P.L., Neely, G.G., et al. (2019a). High Dietary Sugar Reshapes Sweet Taste to Promote Feeding Behavior in Drosophila melanogaster. Cell Rep. 27, 1675–1685.e7.

May, C.E., Rosander, J., Gottfried, J., Dennis, E., and Dus, M. (2019b). Dietary sugar inhibits satiation by decreasing the central processing of sweet taste. bioRxiv.

McCluskey, L.P., He, L., Dong, G., and Harris, R. (2020). Chronic exposure to liquid sucrose and dry sucrose diet have differential effects on peripheral taste responses in female rats. Appetite 145, 104499.

Müller, J., and Kassis, J.A. (2006). Polycomb response elements and targeting of Polycomb group proteins in Drosophila. Curr. Opin. Genet. Dev. 16, 476–484.

Musselman, L.P., and Kühnlein, R.P. (2018). Drosophila as a model to study obesity and metabolic disease. J. Exp. Biol. 221.

Na, J., Musselman, L.P., Pendse, J., Baranski, T.J., Bodmer, R., Ocorr, K., and Cagan, R. (2013). A Drosophila Model of High Sugar Diet-Induced Cardiomyopathy. PLoS Genetics 9, e1003175.

Nanney, D.L. (1958). EPIGENETIC CONTROL SYSTEMS. Proc. Natl. Acad. Sci. U. S. A. 44, 712–717.

Nestler, E.J., and Lüscher, C. (2019). The Molecular Basis of Drug Addiction: Linking Epigenetic to Synaptic and Circuit Mechanisms. Neuron 102, 48–59.

Neumann, C.J., and Cohen, S.M. (1998). Boundary formation in Drosophila wing: Notch activity attenuated by the POU protein Nubbin. Science 281, 409–413.

Parrish, J.Z., Kim, M.D., Jan, L.Y., and Jan, Y.N. (2006). Genome-wide analyses identify transcription factors required for proper morphogenesis of Drosophila sensory neuron dendrites.
Genes Dev. 20, 820–835.

Parrish, J.Z., Emoto, K., Jan, L.Y., and Jan, Y.N. (2007). Polycomb genes interact with the tumor suppressor genes hippo and warts in the maintenance of Drosophila sensory neuron dendrites. Genes Dev. 21, 956–972.

Parrish, J.Z., Kim, C.C., Tang, L., Bergquist, S., Wang, T., Derisi, J.L., Jan, L.Y., Jan, Y.N., and Davis, G.W. (2014). Krüppel mediates the selective rebalancing of ion channel expression. Neuron 82, 537–544.

Qi, W., Zhao, K., Gu, J., Huang, Y., Wang, Y., Zhang, H., Zhang, M., Zhang, J., Yu, Z., Li, L., et al. (2017). An allosteric PRC2 inhibitor targeting the H3K27me3 binding pocket of EED. Nat. Chem. Biol. 13, 381–388.

Quinlan, A.R., and Hall, I.M. (2010). BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics 26, 841–842.

Ragunathan, K., Jih, G., and Moazed, D. (2015). Epigenetics. Epigenetic inheritance uncoupled from sequence-specific recruitment. Science 348, 1258699.

Ramírez, F., Ryan, D.P., Grüning, B., Bhardwaj, V., Kilpert, F., Richter, A.S., Heyne, S., Dündar, F., and Manke, T. (2016). deepTools2: a next generation web server for deep-sequencing data analysis. Nucleic Acids Res. 44, W160–W165.

Sartor, F., Donaldson, L.F., Markland, D.A., Loveday, H., Jackson, M.J., and Kubis, H.-P. (2011). Taste perception and implicit attitude toward sweet related to body mass index and soft drink supplementation. Appetite 57, 237–246.

von Schimmelmann, M., Feinberg, P.A., Sullivan, J.M., Ku, S.M., Badimon, A., Duff, M.K., Wang, Z., Lachmann, A., Dewell, S., Ma'ayan, A., et al. (2016). Polycomb repressive complex 2 (PRC2) silences genes responsible for neurodegeneration. Nat. Neurosci. 19, 1321–1330.

Scott, K. (2018). Gustatory Processing in Drosophila melanogaster. Annu. Rev. Entomol. 63, 15–30.

Shannon, P., Markiel, A., Ozier, O., Baliga, N.S., Wang, J.T., Ramage, D., Amin, N., Schwikowski, B., and Ideker, T. (2003). Cytoscape: a software environment for integrated models of biomolecular interaction networks. Genome Res. 13, 2498–2504.

Shiraiwa, T., and Carlson, J.R. (2007). Proboscis extension response (PER) assay in Drosophila. J. Vis. Exp. 193.

Southall, T.D., Gold, K.S., Egger, B., Davidson, C.M., Caygill, E.E., Marshall, O.J., and Brand, A.H. (2013). Cell-type-specific profiling of gene expression and chromatin binding without cell isolation: assaying RNA Pol II occupancy in neural stem cells. Dev. Cell 26, 101–112.

Stark, R., Brown, G., and Others (2011). DiffBind: differential binding analysis of ChIP-Seq peak data. R Package Version 100, 4–3.
van Steensel, B., and Henikoff, S. (2000). Identification of in vivo DNA targets of chromatin proteins using tethered dam methyltransferase. Nat. Biotechnol. 18, 424–428.

Stewart, J.E., and Keast, R.S.J. (2012). Recent fat intake modulates fat taste sensitivity in lean and overweight subjects. Int. J. Obes. 36, 834–842.

Sweatt, J.D. (2013). The emerging field of neuroepigenetics. Neuron 80, 624–632.

Szklarczyk, D., Gable, A.L., Lyon, D., Junge, A., Wyder, S., Huerta-Cepas, J., Simonovic, M., Doncheva, N.T., Morris, J.H., Bork, P., et al. (2019). STRING v11: protein–protein association networks with increased coverage, supporting functional discovery in genome-wide experimental datasets. Nucleic Acids Res. 47, D607–D613.

Tennesen, J.M., Barry, W.E., Cox, J., and Thummel, C.S. (2014). Methods for studying metabolism in Drosophila. Methods 68, 105–115.

Thurmond, J., Goodman, J.L., Strelets, V.B., Attrill, H., Gramates, L.S., Marygold, S.J., Matthews, B.B., Millburn, G., Antonazzo, G., Trovisco, V., et al. (2019). FlyBase 2.0: the next generation. Nucleic Acids Res. 47, D759–D765.

Tsankova, N.M., Berton, O., Renthal, W., Kumar, A., Neve, R.L., and Nestler, E.J. (2006). Sustained hippocampal chromatin regulation in a mouse model of depression and antidepressant action. Nat. Neurosci. 9, 519–525.

Ueno, K., and Kidokoro, Y. (2008). Adenylyl cyclase encoded by AC78C participates in sugar perception in Drosophila melanogaster. Eur. J. Neurosci. 28, 1956–1966.

Vorbrüggen, G., Constien, R., Zilian, O., Wimmer, E.A., Dowe, G., Taubert, H., Noll, M., and Jäckle, H. (1997). Embryonic expression and characterization of a Ptx1 homolog in Drosophila. Mech. Dev. 68, 139–147.

Wang, Q.-P., Lin, Y.Q., Zhang, L., Wilson, Y.A., Oyston, L.J., Cotterell, J., Qi, Y., Khuong, T.M., Bakhshi, N., Planchenault, Y., et al. (2016). Sucralose Promotes Food Intake through NPY and a Neuronal Fasting Response. Cell Metab. 24, 75–90.

Weiss, M.S., Hajnal, A., Czaja, K., and Di Lorenzo, P.M. (2019). Taste Responses in the Nucleus of the Solitary Tract of Awake Obese Rats Are Blunted Compared With Those in Lean Rats. Front. Integr. Neurosci. 13, 35.

Wilkinson, L. (2012). Exact and approximate area-proportional circular Venn and Euler diagrams. IEEE Trans. Vis. Comput. Graph. 18, 321–331.

Wise, P.M., Nattrass, L., Flammer, L.J., and Beauchamp, G.K. (2016). Reduced dietary intake of simple sugars alters perceived sweet taste intensity but not perceived pleasantness. Am. J. Clin. Nutr. 103, 50–60.

Zaffran, S., Das, G., and Frasch, M. (2000). The NK-2 homeobox gene scarecrow (scro) is expressed in pharynx, ventral nerve cord and brain of Drosophila embryos. Mech. Dev. 94,
237–241.

Zhang, Y.V., Raghuwanshi, R.P., Shen, W.L., and Montell, C. (2013). Food experience-induced taste desensitization modulated by the Drosophila TRPL channel. Nat. Neurosci. 16, 1468–1476.
Figure legends

Figure 1 PRC2.1 modulates sweet taste in response to the food environment.

(A) Schematic of *Drosophila melanogaster* PRC2 core complex consisting of Enhancer of zeste (E(z)), Suppressor of zeste (Su(z)12), extra sex combs (esc) and the accessory protein Polycomb-like (Pcl). (B-G) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males (B) w1118CS (gray), E(z)c249/+ (dark red), Su(z)12c253/+ (fuschia), and escc289/+ (red) flies on a control (circle) or sugar (square) diet for 7 days. n = 34-68; (C) w1118cs flies on a control (circle) or sugar (square) diet supplemented with vehicle (10% DMSO) or 8uM EEDi for 7 days. n = 32-43; (D) w1118cs (gray), Pclp429/+ (pink) and PclCI1/+ (pink) flies on a control (circle) or sugar (square) diet for 7 days. n = 36-82; (E) Gr5a>PclRNAi-1 (pink), and parental transgenic controls (gray, crossed to w1118cs) flies on a sugar diet for 7 days. n = 42-63; (F) Gr5a>Pcl (pink), and parental transgenic controls (gray, crossed to w1118cs) flies on a control diet for 7 days. n = 36-61; (G) Gr5a>Pcl flies on a control diet supplemented with vehicle (10% DMSO) or 8uM EEDi for 7 days. n = 30-35. Data are shown as mean ± SEM, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to control diet within genotype groups. **** p < 0.0001, *** p < 0.001, ** p < 0.01,*p < 0.05 for all panels in this figure.

Figure 2 Pcl mutant flies have normal sensory responses and are resistant to diet-induced obesity.

(A) Schematic showing the setup for in vivo calcium imaging: the fly proboscis is stimulated with 20% sucrose while recording from the subesophageal zone (SEZ) which contains the presynaptic terminals of the sweet taste neurons (shown here labeled with
synaptagmin::GFP). (B) Average ΔF/F₀ calcium response traces (left panel) and peak % ΔF/F₀ (right panel) responses to sucrose stimulation of the proboscis (arrow) in age-matched male Gr64f>GCaMP6s-Bruchpilot-mCherry;Pcl<sup>E429</sup> flies fed a control (gray, circle) or sugar (pink, square) diet for 7 days, n = 32-34. (C) Quantification of GFP-labeled cells in the labella of Gr64f;CD8-GFP flies crossed to w<sup>1118cs</sup> (as control, dark gray) or Gr64f;CD8-GFP>Pcl<sup>RNAi</sup> (pink) on a control (circle) or sugar (square) diet for 7 days. n=5-16 flies, Kruskal-Wallis Dunn’s multiple comparisons, comparison to control diet of each genotype, no significance. (D) Triglyceride levels normalized to protein in Gr5a>Pcl<sup>RNAi</sup> (pink) and parental transgenic control flies (dark gray, crossed to w<sup>1118cs</sup>) fed a control (circle) or sugar (square) diet for 7 days. n=8, two-way ANOVA with Sidak’s multiple comparisons test, comparisons to control diet of each genotype. All data are shown as mean ± SEM, ****p < 0.0001, ***p < 0.001, **p < 0.01, and *p < 0.05 for all panels unless indicated.

Figure 3 Pcl chromatin occupancy is redistributed in the high sugar environment.

(A) Schematic of Targeted Dam-ID (TaDa) of Dam::Pcl and Dam (CATaDa) and of the TaDa induction (top panel). Age-matched Gr5a;tubulin-GAL80>UAS-LT3-Dam::Pcl and Gr5a;tubulin-GAL80>UAS-LT3-Dam flies were placed on a control (gray) or sugar (purple) diet for 3 days at 20-21°C, then switched to 28°C between day 3-4 to induce Dam expression (arrow) (bottom panel). (B) CATaDa coverage normalized to 1x dm6 genome (reads per genome coverage) of the three biological replicates from control diet flies within a 5 kb window at the sweet gustatory receptor Gr5a and the bitter gustatory receptor Gr66a. (C) Fold change for the proportion of genes allocated to the five chromatin states (assigned in a color) according to their transcription start site (TSS), normalized to the expected proportion across the whole genome (see methods). (D) Averaged log₂(Dam::Pcl/Dam) tracks for a 10 kb window centered on
predicted PREs on a control (gray) and sugar (purple) diet. (E-F) Averaged CATaDA tracks normalized to 1x dm6 genome for a 10 kb window centered on (E) Pcl peaks and (F) predicted PREs on a control (gray) and sugar (yellow) diet. (G) Box plots represent the median and variance of normalized Dam::Pcl/Dam reads for genes differentially bound by Pcl on a control (gray) and sugar (purple) diet. Genes are placed into groups with higher (group 1) or lower (group 2) binding in the sugar diet environment. (H) GO terms associated with differentially bound genes by Dam::Pcl/Dam identified by iPAGES, text boxes represent the GO term category, metabolism (orange) and Regulatory (lavender) (for the full iPAGE list see Supplemental Figure 4).

Figure 4 PRC2.1 sculpts the transcriptional responses of the Gr5a+ neurons in response to diet. (A) Schematic depiction of the riboTRAP technique used to profile the changes in the Gr5a+ neurons. (B-C) Volcano plots representing changes in gene expression in the Gr5a+ neurons of age matched male Gr5a>UAS-Rpl3-3XFLAG flies on a sugar diet for 3 days (SD3, mint) and 7 days (SD7, teal). n=2-3 replicates per condition (10,000 cells per replicate). Non-significant genes are shown in black and genes with q<0.1 (Wald test) are in mint or teal for SD3 and SD7 respectively, comparison is to the control diet group. (D) Venn diagram of differentially expressed genes (DEGs) at SD3/CD (mint), SD7/CD (teal), and the overlap between SD3/SD7 (Wald test, q<0.1). The heatmaps show log2 fold changes (l2 fc) for DEGs in each condition specified in the venn diagram (left columns, SD3, SD7, and SD3+SD7, all comparisons to control diet) and Pcl$^{429}$ mutant flies (right column, Pcl$^{429}$ SD7 comparison to control diet); l2 fc range from purple (high) to gold (low). (E) Normalized read counts (Tpm+1) from TRAP for Pcl bound (pink) and not bound genes (gray) on a control diet. Box plots represent the median and
variance, \( p\text{-value} = 3.196 \times 10^{-6} \), two-tailed t-test. (F) \( \log_2 \) fold changes for Scarecrow (Scro), Paired-like homeobox domain 1 (Ptx1), caudal (cad), GATAe, and nubbin (nub) in control SD7/CD and SD3/CD flies, and \( Pcf^{429} \) SD7/CD mutant flies. \( \log_2 \text{fc} \) range from purple (high) to gold (low).

**Figure 5 PRC2.1 represses a transcriptional program required for sweet taste.**

(A) \( \log_2 \) [Dam::Pcl/Dam] coverage tracks of flies on a control (black) and sugar (purple) diet within a 50 kb window at the \( cad \), \( Pttx1 \), GATAe, nub, and Scro genes visualized with the UCSD genome browser. The three biological replicates for each condition are superimposed. Turquoise traces are SD/CD fold changes. Peaks are shown as black boxes below respective traces (q<0.01) and genes are shown in dense format to include all potential isoforms. (B) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) in age-matched males of \( Gr5a>cad^{RNAi} \) (gold), \( Gr5a>Ptx1^{RNAi} \) (blue), \( Gr5a>GATAe^{RNAi} \) (green), \( Gr5a>nub^{RNAi} \) (lavender), \( Gr5a>Scro \) (salmon), and parental transgenic control (gray, crossed to \( w^{1118} \)) flies on a control diet for 7 days. \( n = 30-54 \), Kruskal-Wallis Dunn’s multiple comparisons, comparisons to both transgenic controls. All data shown as mean ± SEM, ****p < 0.0001, ***p < 0.001, **p < 0.01, and *p < 0.05. (C) Box plots represent the median and variance of the Log\(_2\) fold changes (\( \log_2 \text{fc} \)) for candidate gene targets of \( cad \), \( Pttx1 \), GATAe, nub, and Scro (for threshold cutoff score see Supplemental Figure 10) in control flies at SD7 (teal), and \( Pcf^{429} \) mutants at SD7 (pink). (D) Venn diagram representation of the overlap of the candidate gene targets of \( cad \) (gold), \( Pttx1 \) (blue), GATAe (green), nub (lavender), and Scro (salmon). (E) A transcriptional loop between \( cad \) (gold), \( Pttx1 \) (blue), GATAe (green), nub (lavender), and Scro (salmon) mediated by Pol. Green arrows represent activation and red bars represent inhibition. GO terms associated with each transcriptional regulator identified by iPAGE
are listed. Text boxes represent the GO term category, metabolism (orange), regulatory (lavender) and neural/signal (blue) (for the full iPAGE list see Supplemental Figure 11-13).

**Figure 6** The persistent phenotypic memory of the food environment is dependent on PRC2.1.

(A) Diagram representing dietary manipulations, CD, control diet (gray), SD, sugar diet (purple), and > represents the switch in diet. (B) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males w1118^{2a} flies on a control (circle), control to sugar (triangle), and sugar to control (diamond) diet as shown in (A) n =57-64, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to control diet. All data shown as mean ± SEM, ****p < 0.0001, ***p < 0.001, **p < 0.01, and *p < 0.05 in this figure. (C) Volcano plot with the changes in gene expression in the Gr5a+ neurons of age-matched male Gr5a>UAS-Rpl3-3XFLAG flies on a sugar to control (SD>CD, green) diet compared to the control diet group (Wald test, q<0.1, ns is non-significant). n=2 replicates per condition (~10,000 cells per replicate). (D) Heatmap showing how the differentially expressed genes identified in Figure 5C are changing in the SD>CD condition (310/658, 48% Wald test, q<0.1) compared to the other conditions SD3, SD7, and Pcf^{+/2}SD7 (column headings, all comparisons to control diet). Log_{2} fold changes for these genes were determined by comparing each group to its own control (see methods). I_{2}fc range from purple (high) to gold (low). (E) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males w1118^{2a} flies on a control (circle), control to sugar (triangle), and sugar to control (diamond, green) diet for 14 days. EEDi was supplemented to the diets at the day 7 transition (see diagram in figure). n =34-46, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to control diet. (F) Model of molecular changes in the Gr5a+ neurons on a control (top panel) and
sugar (bottom panel) diet, showing the redistribution in PRC2.1 binding, and the effects on the transcription of the regulators (green and red arrows), and neural responses to sweet stimuli.
**Supplement figure legends**

**Supplement Figure 1, related to Figure 1**

(A) The fly proboscis contains the sweet sensory neurons soma (green) with dendrites extending into the hair sensilla and their axons projecting onto the subesophageal zone (SEZ, delimited in yellow) of the fly brain. (B) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males of Gr5a> Jarid2RNAi-1 (dark blue), Gr5a>JariδRNAi-2 (light blue), and Gr5a>jingRNAi-1 (orange) and parental transgenic controls (gray, crossed to w1118 CS) flies on a sugar diet for 7 days. n = 25-34, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to transgenic controls. (C) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males of w1118 CS (gray), ph-d (401j+), ph-p (602j+), Psc (27j+), and Pc 1/2 flies on a control (circle) or sugar (square) diet for 7 days. n = 30-42, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to control diet within each genotype group. (D) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males of w1118 CS (gray), phol (81A) flies on a control (circle) or sugar (square) diet for 7 days. n = 32-34, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to control diet within each genotype group. (E) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males of Gr5a>Sfmbt RNAi (dark pink) and parental transgenic controls (gray, crossed to w1118 CS) flies on a sugar diet for 7 days. n = 26-46, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to transgenic controls. All data shown as mean ± SEM, ****p < 0.0001, ***p < 0.001, **p < 0.01, and *p < 0.05 for all panels unless indicated.

**Supplement Figure 2, Related to Figure 1**
(A) Fold change of Pcl mRNA levels in the heads of flies with and without pan-neuronal (nsyb-GAL4) Pcl knockdown with two independent RNAi lines (dark gray, plum, and pink respectively) measured by qPCR and normalized to the gene Rp49. n=6, Kruskal-Wallis Dunn’s multiple comparisons, compared to GAL4 transgenic control (crossed to w1118^b). (B) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males of Gr5a>PclRNAi-2 and parental transgenic controls (gray, crossed to w1118^b) flies on a sugar diet for 7 days. n = 42-51, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to transgenic controls. (C) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males of Gr5a>PclRNAi-1 and parental transgenic controls (gray, crossed to w1118^b) flies on a control diet for 7 days. n =30-35, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to transgenic controls. (D) Quantification of the number of sweet taste GFP-labeled cells in the labella of Gr64f;CD8-GFP flies crossed to w1118^b (as control, gray) or Gr64f;CD8-GFP>Pcl (pink) on a control (circle) or sugar (square) diet for 7 days. n=5-16 probosces, no significance, Kruskal-Wallis Dunn’s multiple comparisons, comparison to control diet of each genotype. (E) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males of Gr5a;tubulin-GAL80^+>PclRNAi-1, Gr5a;tubulin-GAL80^+>Pcl, and parental transgenic control (gray, crossed to w1118^b) flies on a control (circle, right) or sugar (square, left) diet for 7 days. n = 34-49, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to transgenic control. All data shown as mean ± SEM, ****p < 0.0001, ***p < 0.001, **p < 0.01, and *p < 0.05 for all panels unless indicated.

Supplement Figure 3, related to Figure 3
(A) Principal component analysis of normalized log₂(Dam::Pcl/Dam) flies on a control (CD, gray) or sugar (SD, purple) diet. (B) The three biological replicates of log₂(Dam::Pcl/Dam) on a control (CD, gray) or sugar (SD, purple) diet are shown, with high Pearson correlation coefficients within each dietary condition, and low correlation coefficients between dietary conditions. Scale shows degree of correlation. (C-D) Left panel: overlap of log₂(Dam::Pcl/Dam) peaks between a control (gray) and sugar (purple) diet (find_peaks, q<0.01). Right panel: overlap of CATaDA peaks between a control (gray) and sugar (yellow) diet (MACS2, q<0.05). (E-F) Log₂(Dam::Pcl/Dam) coverage at Ultrabithorax (Ubx), abdominal A (abd-A), Abdominal B (Abd-B), polyhomeotic distal (ph-d) and polyhomeotic proximal (ph-p) (Top panel, left), and Tat interactive protein 60 (Tip60) and Suppressor of variegation 3-9 (Su(var)3-9) (Top panel, right). Bottom panel: CATaDA coverage of the three biological replicates at the loci mentioned above. (G) Volcano plot showing the differential binding of Dam::Pcl on a sugar diet compared to a control diet, (Wald test, q<0.05, purple), non significant genes peaks are shown in black (ns).

**Supplement Figure 4, related to Figure 3**

iPAGE identification of pathways depleted (left) or enriched (right) compared to background gene list (middle) in the differentially bound peaks by Dam::Pcl flies on a sugar diet compared to a control diet. Scale represents over-representation (orange) or under-representation (purple) of genes within a specific bin for the corresponding GO term. Black boxes represent q<0.05.

**Supplement Figure 5, related to Figure 3**

(A) Principal component analysis of age matched male Gr5a>UAS-Rpl3-3XFLAG flies on a control diet (CD, gray), sugar diet for three days (SD3, mint), and sugar diet for seven days (SD7, teal). (B) Normalized read counts for Gustatory receptor Gr66a (bitter), Gustatory
receptor Gr64f, Gustatory receptor Gr5a, and Gustatory receptor Gr64a (all sweet) from the input fraction (dark green box) and Gr5a+ fraction (light green box) specific conditions. Both samples also include the dietary conditions (not specified in figure), hues of purple from light to dark represent low to high expression. (C) Principal component analysis of age matched male Gr5a;Pcr\textsuperscript{kr29}>UAS-Rpl3-3XFLAG flies on a control diet (CD, gray) and sugar diet for seven days (SD7, pink). (D) Volcano plot with changes in gene expression in the Gr5a+ neurons of age matched male Gr5a;Pcr\textsuperscript{kr29}>UAS-Rpl3-3XFLAG flies on a SD for 7 days (Pcr\textsuperscript{kr29}SD7, pink) compared to the control diet condition in the same genotype (Wald test, q<0.1), n=3 replicates per condition (~10,000 cells per replicate). (E) Normalized reads (Tpm+1) from TRAP for genes bound by Pcl (x-axis) and normalized to Dam::Pcl/Dam reads at these genes (y-axis) on a control diet. Scale represents gene density (light to dark blue). (F) Fold change of Dam::Pcl binding for genes differentially bound on sugar diet compared to control diet (y-axis) (q<0.05) and their respective differentially expressed log\textsubscript{2} fold changes on a sugar diet compared to control diet (x-axis) (q<0.1). Genes are shown as circles and colored based on GO term category, metabolism (orange) and regulatory (lavender).

**Supplement Figure 6, related to Figure 4**

iPAGE identification of pathways enriched in age-matched male Gr5a>UAS-Rpl3-3XFLAG flies on a sugar diet for three days compared to a control diet. Bins (top) show the range of log\textsubscript{2} fold changes for genes within their corresponding GO terms. Scale represents over-representation (orange) or under-representation (purple) of genes within a specific bin for the corresponding GO term. Black boxes represent q<0.05.

**Supplement Figure 7, related to Figure 4**
iPAGE identification of pathways enriched in age-matched male *Gr5a>*UAS-Rpl3-3XFLAG flies on a sugar diet for seven days compared to a control diet. Bins (top) show the range of log$_2$ fold changes for genes within their corresponding GO terms. Scale represents over-representation (orange) or under-representation (purple) of genes within a specific bin for the corresponding GO term. Black boxes represent *q*<0.05.

**Supplement Figure 8, related to Figure 4**

iPAGE identification of pathways enriched in age-matched male

*Gr5a>*Pd$^{K429}$>*UAS-Rpl3-3XFLAG flies on a sugar diet for seven days compared to a control diet. Bins (top) show the range of log$_2$ fold changes for genes within their corresponding GO terms. Scale represents over-representation (orange) or under-representation (purple) of genes within a specific bin for the corresponding GO term. Black boxes represent *q*<0.05.

**Supplement Figure 9, related to Figure 5**

(A) Normalized read counts for *Scro, cad, nub, GATAe*, and *Ptx1* in the input fraction (dark green box) and the *Gr5a*+ fraction (light green box) from *Gr5a>*UAS-Rpl3-3XFLAG age-matched male flies on a control diet; hues of purple from light to dark show low to high expression. (B) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males of *Gr5a>*cad (gold), *Gr5a>*Ptx1 (blue), *Gr5a>*nub (lavender), *Gr5a>*Scro$^{RNAi}$ (coral), and parental transgenic control (gray, crossed to w$^{1118cs}$) flies on a sugar diet for 7 days. *n* = 31-51, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to transgenic controls.

**Supplement Figure 10, related to Figure 5**
(A) Schematic of how the targets of the 5 transcription factors in Figure 5 were identified: the DNA binding motifs for each transcription factor were scanned along every base pair of the *D. melanogaster* genome using FIMO, and the basepair-wise binding results converted to robust z-scores. (B-F) Threshold scores used to identify the regulons of each transcription factor. Y-axis represents the averaged log₂ fold change of the predicted targets based on the cutoff score, respectively.

**Supplement Figure 11, related to Figure 5**
iPAGE identification of pathways over-represented (orange) or under-represented (purple) in the predicted gene targets of *GATAe* compared to all other genes that do not fit the cutoff thresholds from Supplemental Figure 10 (non-targets). Black boxes represent *q*<0.05.

**Supplement Figure 12, related to Figure 5**
iPAGE identification of pathways over-represented (orange) or under-represented (purple) in the predicted gene targets of *cad*, and *Scro* independent of each other to all other genes that do not fit the cutoff thresholds from Supplemental Figure 10 (non-targets). Black boxes represent *q*<0.05.

**Supplement Figure 13, related to Figure 6**
iPAGE identification of pathways over-represented (orange) or under-represented (purple) in the predicted gene targets of *nub/pdm* and *Pttx1* independent of each other to all other genes that do not fit the cutoff thresholds from Supplemental Figure 10 (non-targets). Black boxes represent *q*<0.05.
Supplement Figure 14, related to Figure 6

(A) Functional network created with STRING featuring PRC2.1, cad, Ptx1, GATAe, nub, Scro, and their predicted targets from Figure 5C that can be classified into the GO term category neural/signaling. Each node represents a gene. Edges indicate functional protein-protein interactions, not direct interactions. Edges represented with dashed lines are interactions identified in this study. (B) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males of w1118cs, Arc1esm18, and Arc1esm13 flies on a control diet for 7 days. n = 38-45, Kruskal-Wallis Dunn’s multiple comparisons, comparisons to transgenic controls. (B) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males of Gr5a>Arc1RNAi (turquoise), and parental transgenic control (gray, crossed to w1118cs) flies on a control diet for 7 days. n = 28-47, Kruskal-Wallis Dunn’s multiple comparisons, comparison to control diet. (C) Taste responses (y axis) to stimulation of the labellum with 30%, 10% and 5% sucrose (x axis) of age-matched males of Gr5a>ACXDRNAI-1 (dark purple) and Gr5a>ACXDRNAI-2 (light purple), and parental transgenic controls (gray, crossed to w1118cs) flies on a control diet for 7 days. n = 28-47, Kruskal-Wallis Dunn’s multiple comparisons, comparison to control diet. (D) Triglyceride levels normalized to protein in w1118cs flies fed a control (gray, circle) or sugar (gray, square) diet for 14 days, and flies fed a SD>CD reversal diet for a total of 14 days (7 days in each diet). n=16, two-way ANOVA with Sidak’s multiple comparisons test, comparisons to control diet of each genotype. All data are shown as mean ± SEM, ****p < 0.0001, ***p < 0.001, **p < 0.01, and *p < 0.05 for all panels unless indicated.

Supplement Figure 15, related to Figure 6
iPAGE identification of pathways enriched in age-matched male Gr5a>UAS-Rpl3-3XFLAG flies on a SD>CD compared to flies on a CD>CD. Bins (top) show the range of log₂ fold changes for genes within their corresponding GO terms. Scale represents over-representation (orange) or under-representation (purple) of genes within a specific bin for the corresponding GO term. Black boxes represent q<0.05.

Supplementary Files

Supplementary file 1:
PRE prediction calls for the dmel6.08 genome. Relating to Figure 3.

Supplementary file 2:
Differential binding analysis of Dam::Pcl/Dam on a sugar diet compared to control diet.

Supplementary file 3:
Differential expression analysis of riboTRAP experiments for sugar diet 3 (SD3), sugar diet 7 (SD7), Pcl<sup>429</sup> SD7 (all comparisons to control diet) and CD>CD and SD>CD.

Supplementary file 4:
Transcription factor regulatory target calls relating to Figures 5.

Supplementary file 5:
STRING interaction network relating to Supplement Figure 14.
Supplementary file 6:

Gene table for heatmap relating to Figure 6D.
Figure 2

Vaziri et al.

A

Sweet taste neuron axons
20% sucrose

B

PCk429

Sucrose

Peak % ΔF/F₀

C

Gr5a-GAL4  Gr5a>Pcl₉₄₉

# of GFP⁺ cells

CD  SD

D

Gr5a-GAL4  Gr5a>Pcl₁₄₄₉

Triglycerides/Protein

CD  SD
**Vaziri et al.**

**Figure 3**

A. Diagram showing the interaction between Dam::Pcl and DNA.

B. Gene count fold change graphs for Rep 1, Rep 2, and Rep 3.

C. Bar graph showing gene count fold change with color-coded bars.

D. Log2 (Dam::Pcl/Dam) SD/CD GO enrichment for Metabolic processes, Regulatory, Transcription repressor activity, Neuroblast fate commitment, Axon target recognition, and Proboscis development.
Vaziri et al.
Figure 6

A

CD > CD
CD > SD
SD > CD

Days on diet

B

CD > CD
CD > SD
SD > CD

PER

Log_{10}(q-value)

Log_{2}(fold change)

C

ns
q<0.1

skew=-2.06

D

SD > CD
SD7
SD3
Pctl^{429} SD7

E

CD > CD + EEDi
CD > SD + EEDi
SD > CD + EEDi

PER

Log_{10}(q-value)

Log_{2}(fold change)

F

Control Diet
Sugar Diet

Scro
GATAe
cad
Pt1
Ptx1
nub

310 genes
A

B

C

D

E

Vaziri et al.
Supplement 2
transcriptional repressor activity, RNA polymerase II core promoter proximal region, GO:0001078
hydrolase activity, acting on carbon-nitrogen (but not peptide) bonds, in linear amides, GO:0016811
arylamine N-acetyltransferase activity, GO:0004060
peptide disulfide oxidoreductase activity, GO:0015037
response to water, GO:0009415
protein tyrosine phosphatase activity, metal-dependent, GO:0030946
pharynx development, GO:0060465
positive regulation of cell cycle process, GO:00090068
calcium:sodium antiporter activity, GO:0005432
elongation of arista core, GO:0035015
RNA polymerase II transcription factor activity, sequence-specific DNA binding, GO:0000981
axon target recognition, GO:0007412
spiracle morphogenesis, open tracheal system, GO:0035277
reproductive structure development, GO:0048608
proboscis development, GO:0048728
regulation of nervous system development, GO:0051960
N-acetylmuramoyl-L-alanine amidase activity, GO:0008745
glutamate dehydrogenase [NAD(P)+] activity, GO:0004353
negative regulation of fibroblast growth factor receptor signaling pathway, GO:0040037
positive regulation of female receptivity, GO:0045925
co-SMAD binding, GO:0070410
sodium channel regulator activity, GO:0017080
serine-type endopeptidase inhibitor activity, GO:0004867
transcription factor activity, RNA polymerase II distal enhancer binding, GO:0003705
larval feeding behavior, GO:0030536
negative regulation of antibacterial peptide biosynthetic process, GO:0002809
male courtship behavior, orientation prior to leg tapping and wing vibration, GO:0016543
intercalary heterochromatin, GO:0005725
neuroblast fate commitment, GO:0014017
nucleolus, GO:0005730
G-protein coupled amine receptor activity, GO:0008227
cGMP biosynthetic process, GO:0006182
acetylcholine-activated cation-selective channel activity, GO:0004889
voltage-gated calcium channel complex, GO:0005891
snRNA pseudouridylation guide activity, GO:0030565
voltage-gated potassium channel complex, GO:0008076
gamma-tubulin binding, GO:0043015
positive regulation of glial cell proliferation, GO:0060252
G-protein coupled receptor activity, GO:0004930
sodium channel regulator activity, GO:0017080
sensory perception of chemical stimulus, GO:0007606
deactivation of rhodopsin mediated signaling, GO:0016059
histone H3 acetylation, GO:0043966
dendrite, GO:0030425
vesicle-mediated transport, GO:0016192
neuromuscular junction, GO:0031594
kinesin binding, GO:0019894
precatalytic spliceosome, GO:0071011
guanyl-nucleotide exchange factor activity, GO:0005085
GTPase activator activity, GO:0005096
circadian rhythm, GO:0007623
DNA repair, GO:0006281
mitotic sister chromatid separation, GO:0051306
neuron projection morphogenesis, GO:0048812
asymmetric neuroblast division, GO:0055059
RNA processing, GO:0006396
compound eye development, GO:0048749
protein complex, GO:0043234
muscle organ development, GO:0007517
TORC1 signaling, GO:0038202
transcription corepressor activity, GO:0003714
positive regulation of JNK cascade, GO:0046330
nuclear pore organization, GO:0006999
nucleosome, GO:0000786
cytosolic ribosome, GO:0022626
regulation of protein kinase activity, GO:0045859
chitin-based cuticle development, GO:0040003
mitochondrial large ribosomal subunit, GO:0005762
NADH dehydrogenase activity, GO:0003954
ligand-gated ion channel activity, GO:0015276
endomembrane system, GO:0012505
pyridoxal phosphate binding, GO:0030170
proteasome-mediated ubiquitin-dependent process, GO:0043161
glutathione transferase activity, GO:0004364
metabolic process, GO:0008152
plasma membrane proton-transporting V-type ATPase complex, GO:0033181
carbohydrate metabolic process, GO:0005975
UDP-glucose metabolic process, GO:0006011
transporter activity, GO:0005215
glycolytic process, GO:0006096
serine-type endopeptidase inhibitor activity, GO:0004867
dynein complex, GO:0030286
transmembrane transporter activity, GO:0022857
aminopeptidase activity, GO:0004177
secondary active organic cation transporter activity, GO:0008513
iron ion binding, GO:0005050
alkaline phosphatase activity, GO:0004035
defense response to Gram-positive bacterium, GO:0050830
metallocarboxypeptidase activity, GO:0004181
ketohexokinase activity, GO:0004454
hexose metabolic process, GO:0019318
alpha-mannosidase activity, GO:0004559
lipase activity, GO:0016298
sperm storage, GO:0046693
glucose transmembrane transporter activity, GO:0005355
larval serum protein complex, GO:0005616
A B

Input
Gr5a+

| log₂(TPM+1) | 0 | 1 | 2 | 3 |
|-------------|---|---|---|---|
| nub         |   |   |   |   |
| Scro        |   |   |   |   |
| Ptx1        |   |   |   |   |
| cad         |   |   |   |   |
| GATAe       |   |   |   |   |

**Gr5a-GAL4/+ UAS/+**

**Gr5a+/+**

**PER**

30 10 5
[ΣSucrose]
| Background | Targets |
|------------|---------|
| defense response to Gram-positive bacterium, GO:0050830 | defense response to bacterium, GO:0009617 |
| response to bacterium, GO:0009617 | Rab protein signal transduction, GO:0032482 |
| peptidoglycan binding, GO:0042833 | ventral cord development, GO:0007419 |
| signal transduction, GO:0007165 | mitotic spindle assembly, GO:0090307 |
| phosphoprotein phosphatase activity, GO:004721 | defense response, GO:0006952 |
| presynaptic membrane, GO:0042734 | perinuclear region of cytoplasm, GO:0048471 |
| heme binding, GO:0020307 | transmembrane signaling receptor activity, GO:0004888 |
| positive regulation of cell proliferation, GO:0006284 | phosphoprotein phosphatase activity, GO:0004721 |
| phosphorylation, GO:0016311 | regulation of synaptic growth at neuromuscular junction, GO:0008582 |
| defense response to fungus, GO:0005832 | serine-type endopeptidase inhibitor activity, GO:0004867 |
| Golgi apparatus, GO:0005794 | iron ion binding, GO:0006952 |
| iron ion binding, GO:0005506 | receptor activity, GO:0004872 |
| metabolic process, GO:0008152 | regulation of cell shape, GO:0006284 |
| regulation of cell shape, GO:0008360 | apical plasma membrane, GO:0016324 |
| cell adhesion, GO:0007155 | border follicle cell migration, GO:0007298 |
| chitin binding, GO:0006801 | positive regulation of GTPase activity, GO:0043547 |
| RNA binding, GO:0003723 | carbohydrate binding, GO:0003024 |
| regulation of cell cycle, GO:0001700 | glutathione metabolic process, GO:0006749 |
| fatty acid beta-oxidation, GO:0006635 | ATC-binding cassette (ABC) transporter complex, GO:0043190 |
| synaptic vesicle, GO:0008021 | ATP-binding cassette (ABC) transporter complex, GO:0043190 |
| vesicle-mediated transport, GO:0016192 | transcription factor binding, GO:0008134 |
| transcription factor binding, GO:0008134 | carbohydrate metabolic process, GO:0005975 |
| carbohydrate metabolic process, GO:0005975 | ATP-binding cassette (ABC) transporter complex, GO:0043190 |
| AP-binding cassette (ABC) transporter complex, GO:0043190 | lipase activity, GO:0016298 |
| lipase activity, GO:0016298 | nucleotide binding, GO:000166 |
| nucleotide binding, GO:000166 | pyridoxal phosphate binding, GO:000170 |
| pyridoxal phosphate binding, GO:000170 | recycling endosome, GO:0005037 |
| recycling endosome, GO:0005037 | subapical complex, GO:0035003 |
| subapical complex, GO:0035003 | protein kinase activity, GO:0004672 |
| protein kinase activity, GO:0004672 | protein kinase CK2 complex, GO:0005956 |
| protein kinase CK2 complex, GO:0005956 | ribosome, GO:0005840 |
| ribosome, GO:0005840 | intracellular membrane-bounded organelle, GO:0043231 |
| intracellular membrane-bounded organelle, GO:0043231 | endocytosis, GO:0008897 |
| endocytosis, GO:0008897 | endocytic recycling, GO:0032456 |
| endocytic recycling, GO:0032456 | neuron projection morphogenesis, GO:0048812 |
| neuron projection morphogenesis, GO:0048812 | mitotic cell cycle, GO:0000278 |
| mitotic cell cycle, GO:0000278 | neurotransmitter secretion, GO:0007269 |
| neurotransmitter secretion, GO:0007269 | nucleosome, GO:0000786 |
| nucleosome, GO:0000786 | male meiosis, GO:0007140 |
| male meiosis, GO:0007140 | cytosolic ribosome, GO:0022626 |
| cytosolic ribosome, GO:0022626 | epithelial cell migration, open tracheal system, GO:0007427 |
| epithelial cell migration, open tracheal system, GO:0007427 | chromatin silencing at centromere, GO:0030702 |
| chromatin silencing at centromere, GO:0030702 | embryonic development via the syncytial blastoderm, GO:0001700 |
| embryonic development via the syncytial blastoderm, GO:0001700 | GGC codon-amino acid adaptor activity, GO:0033462 |
| GGC codon-amino acid adaptor activity, GO:0033462 | learning or memory, GO:0007611 |
| learning or memory, GO:0007611 | protein kinase CK2 complex, GO:0005956 |
| protein kinase CK2 complex, GO:0005956 | chromatin assembly or disassembly, GO:0006333 |
| chromatin assembly or disassembly, GO:0006333 | CGA codon-amino acid adaptor activity, GO:0033431 |
| CGA codon-amino acid adaptor activity, GO:0033431 | trachea morphogenesis, GO:00060439 |
| trachea morphogenesis, GO:00060439 | nucleosome assembly, GO:0006334 |
| nucleosome assembly, GO:0006334 | chromatin silencing, GO:0006342 |
| chromatin silencing, GO:0006342 | CCG codon-amino acid adaptor activity, GO:0033421 |
| CCG codon-amino acid adaptor activity, GO:0033421 | auditory behavior, GO:0031223 |
| auditory behavior, GO:0031223 | UGC codon-amino acid adaptor activity, GO:0033414 |
| UGC codon-amino acid adaptor activity, GO:0033414 | heat shock-mediated polypeptide chromosome puffing, GO:0035080 |
| heat shock-mediated polypeptide chromosome puffing, GO:0035080 | morphogenesis of embryonic epithelium, GO:0016331 |
| morphogenesis of embryonic epithelium, GO:0016331 | terminal branching of open tracheal system, GO:0007430 |
| terminal branching of open tracheal system, GO:0007430 | olfactory behavior, GO:0048248 |
| olfactory behavior, GO:0048248 | spermatogenesis, exchange of chromosomal proteins, GO:0035093 |
| spermatogenesis, exchange of chromosomal proteins, GO:0035093 | female meiosis chromosome segregation, GO:0016321 |
| female meiosis chromosome segregation, GO:0016321 | flight behavior, GO:0007629 |
| flight behavior, GO:0007629 | protein kinase regulator activity, GO:0019887 |
| protein kinase regulator activity, GO:0019887 | GATAe |
Vaziri et al.
Supplement 13

**nub**
- tachykinin receptor activity, GO:0004995
- positive regulation of calcium-mediated signaling, GO:0050850
- carboxylic ester hydrolase activity, GO:0052689
- chaeta development, GO:0022416
- extracellular matrix, GO:0031012
- structural constituent of chitin-based larval cuticle, GO:0008010
- carbohydrate binding, GO:0030246
- sterol transport, GO:0015918
- learning, GO:0007612
- muscle tissue development, GO:0060537
- chitin-based cuticle development, GO:0040003
- nucleosome, GO:0000786
- chromatin assembly or disassembly, GO:0006333
- cytosolic ribosome, GO:0022626
- female meiosis chromosome segregation, GO:0016321
- male meiosis, GO:0007140

**Pttx1**
- cell adhesion, GO:0007155
- cAMP-dependent protein kinase complex, GO:0005952
- mitochondrial matrix, GO:0005759
- phosphatidylinositol-3-phosphate binding, GO:0032266
- positive regulation of ERK1 and ERK2 cascade, GO:0070374
- pyridoxal phosphate binding, GO:0030170
- NADH dehydrogenase activity, GO:0030170
- centrosome localization, GO:0051642
- lipid metabolic process, GO:0006629
- autophagy, GO:0000422
- ventral cord development, GO:0007419
- central nervous system morphogenesis, GO:0021551
- carbohydrate metabolic process, GO:0005975
- endoderm development, GO:0007492
- phosphoric diester hydrolase activity, GO:0008081
- unfolded protein binding, GO:0051082
- detection of chemical stimulus involved in sensory perception of smell, GO:0050911
- peroxisome, GO:0005777
- metallopeptidase activity, GO:0008237
- threonine-type endopeptidase activity, GO:0004298
- protein folding, GO:0006457
- late endosome, GO:0005770
- endopeptidase activity, GO:0004175
- iron ion binding, GO:0005506
- nucleosome, GO:0000786
- chromosome organization, GO:0051276
- male meiosis, GO:0007140
- nucleosome assembly, GO:0006334
- trachea morphogenesis, GO:00069439
- chromatin silencing at centromere, GO:0030702
- morphogenesis of embryonic epithelium, GO:0016331
- chromatin assembly or disassembly, GO:0006333
- cytosolic ribosome, GO:0022626
- chromatin silencing, GO:0006342
- guanylate cyclase activity, GO:0004383
- GGC codon-amino acid adaptor activity, GO:0033462
- female meiosis chromosome segregation, GO:0016321
- terminal branching, open tracheal system, GO:0007430
A

B

C

D

E

Vaziri et al.
Supplement 14

Gr5a-GAL4/+ UAS-Arc1^{RNAi}/+

Gr5a>Garc1^{RNAi-1} Gr5a>Garc1^{RNAi-2}

Gr5a-GAL4/+ UAS/+
