Binge feeding-induced Olfactory Cortex Suppression Reduces Satiation

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SUMMARY

Binge eating commonly leads to overeating, but the exact mechanism is unclear. While it is known that experiencing flavor contributes to satiety, the interactions between flavor, feeding rate, and food intake remain unknown. Here, we demonstrate a novel feeding rate-dependent feedback loop between olfactory flavor representation in the anterior olfactory (piriform) cortex (aPC) and food intake. Using miniscopes for *in vivo* calcium imaging in freely foraging mice, we identified specific excitatory neuronal responses to food and water during slow feeding. Switching to binge feeding transformed these specific responses into unspecific global suppression of neuronal activity. Food consumption was predicted by the degree of suppression of neuronal activity in the aPC during binge feeding. Also, food deprivation enhanced neuronal activity suppression. We confirmed the hypothesis that aPC suppression promotes food intake with closed-loop optogenetics experiments. Together, we show that olfactory sensory representation in the aPC reciprocally interacts with consummatory behavior to enhance food intake.

Keywords: Binge eating, flavor, smell, taste, Ca²⁺ imaging, optogenetics

INTRODUCTION

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Eating rapidly in a short period, commonly known as binge eating, reduces satiation, the process leading to satiety. Therefore, eating proceeds beyond homeostatic needs (Scisco et al. 2011; Scisco et al. 2011; Andrade, Greene, and Melanson 2008; Bolhuis et al. 2013; Teo, Dam, and Forde 2020; Hurst and Fukuda 2018; Bolhuis et al. 2014). Satiation is an important feedback signal that reduces food consumption upon food intake. The soup paradox illustrates the intimate relationship between feeding rate and satiation: Energy-dense liquids, like apple juice, offer less long-term satiety compared to their isocaloric solid counterparts, such as apples. Interestingly, however, when the rate of liquid food intake is slowed down, for example, with spoon feeding, its ability to satisfy hunger matches that of solid foods, influencing 24-hour food intake (Mattes 2005). The canonical explanation for the reduction of satiation by binge feeding in comparison to slow feeding is based on the delayed transfer of homeostatic signals from the gastrointestinal

tract to the brain (Slyper 2021; Samakidou et al. 2023; Grove et al. 2022). While visceral satiation based on ingestion and absorption is undisputed, there is also sensory satiation mediated by flavor perception (Chen et al. 2015; Mandelblat-Cerf et al. 2015; Cecil, Francis, and Read 1999; Betley et al. 2015). However, no study has investigated whether alterations in flavor representation during binge feeding can reduce sensory satiation. Such a feedback loop between feeding behavior, appetite, and flavor representation would support the emerging concept that reciprocal interactions between action and perception shape behavior (Buzsáki 2019).

Flavor is a multisensory phenomenon involving multiple interconnected primary sensory areas including the primary olfactory cortex or piriform cortex (PC) representing smell, the gustatory cortex (GC) representing taste, and, to some degree, other sensory cortices for tactile and visual aspects of food (Small 2012; Elliott and Maier 2020). The PC generally represents odor identity in a concentration-invariant population code of activated neurons (Blazing and Franks 2020). Random activation of neuronal subpopulations in the PC can determine conditioned appetitive and aversive behavioral responses (Choi et al. 2011), and the PC population code does not contribute to the signaling of odor valence (Wang et al. 2020). If the PC's role in flavor representation during feeding were limited to identification without affecting specific behavioral outcomes, we would expect a stable representation of flavors in the neuronal response patterns to slow and binge feeding.

RESULTS

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Feeding rate modulates flavor representation in the anterior piriform cortex

To examine how feeding rate affects sensory representation, we established a feeding paradigm combined with calcium (Ca^{2+}) imaging in freely moving mice. We built a liquid food delivery system that feeds mice at two different rates to experimentally induce slow or binge feeding. Mice could voluntarily lick from a lick spout to trigger the delivery of a droplet ($\sim 1.8 \mu L$) of liquid food (Ensure, artificial energy-dense flavored nutrient solution) or water. Over 30 min, mice had access to the spout for a total of 14 minutes, with a pseudorandom order of 2 min long

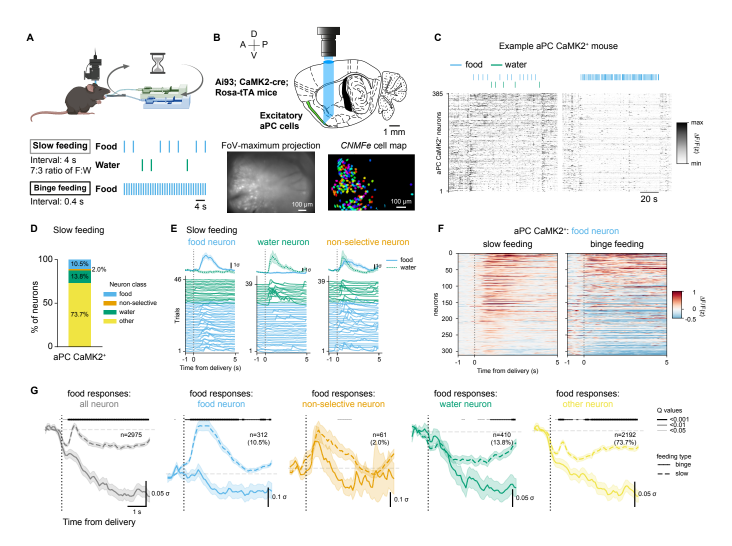


Figure. 1. Feeding rate modulates flavor representations in the aPC.

(A) Miniscope recording and behavioral protocol for slow feeding and binge feeding. (B) Top: Schematics of GRIN lens/Prism (blue shade) implantation in the aPC (green structure). Bottom: Field-of-view of miniscope recordings and extracted cell maps by constrained non-negative matrix factorization (*CN-MFe*). (C) Example Ca²⁺ traces of aPC neurons during slow feeding and binge feeding from one mouse. (D) Percentage of aPC neurons activated during slow feeding by food, water, or non-selective consumption vs. non-responding neurons. (n=2975 cells, 481 slow feeding trials, and 241 binge feeding trials in 8 mice). (E) Trial-average and single trial responses of example aPC food-activated, water-activated, and non-selective neurons upon slow feeding. (F) Trial-averaged responses to food deliveries of individual aPC CaMK2⁺ food-activated neurons upon slow feeding and binge feeding (n=312 cells in 8 mice). (G) Trial-average responses of population and subclasses of aPC CaMK2⁺ neurons upon slow and binge feeding (n is the same as in D). The dashed vertical line indicates the start of the food delivery. The shaded line above denotes the adjusted P-values (Q-values) of each time point, with different line widths representing different values (from thin to thick: Q <0.05, <0.01, <0.001). For (E) and (G) data are shown as mean ± s.e.m.

slow and fast feeding periods. To control the feeding rate, we implemented different refractory periods for the delivery pump: 4 seconds for the slow feeding mode and 0.4 seconds for the binge feeding mode (Fig. 1A). Mice consumed more food and licked at a higher rate during binge feeding (Fig. S1A-D).

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To record feeding-related neuronal activity patterns in the anterior olfactory (piriform) cortex (aPC), we used endomicroscopic lenses (GRIN lenses) attached to prisms combined with a miniscope to image Ca^{2+} transients of excitatory aPC neurons expressing GCaMP6f (Fig. 1B, and S3A-C). We were able to stably record around 140 aPC excitatory neurons per individual imaging session (Fig. S3D, average = 140.40 ± 72.19 neurons).

When switching from slow feeding to binge feeding, we observed a robust suppression of the aPC excitatory neuron population activity (Fig. 1C, Supplementary Video S1). We categorized neurons based on their responses during slow feeding, as this feeding mode provided a larger number of clearly separated individual trials. We interspersed brief water deliveries in the slow feeding mode to differentiate food-specific neurons from non-specific feeding-related neurons. This protocol allowed us to classify neurons pooled over animals and sessions into food-activated (10.5%), water-activated (13.8%), non-selective consumption-activated (neurons responding to both food and water deliveries; 2.0%), and non-responders (73.7%; Fig. 1D, E and S2A, B). Binge feeding-induced aPC suppression was present in food-activated, water-activated, and non-responding neurons but not in non-selective consumption-activated neurons (Fig. 1F, G, S2C-F, statistical analysis of the neuronal activity differences during slow and binge feeding are illustrated with Q-values (unpaired t-test with false discovery rate correction) along the averaged Ca²⁺ traces).

Activity in the PC represents odor identity through a distributed population code and transmits information about odor identity and concentration to downstream targets (Miura, Mainen, and Uchida 2012; Stettler and Axel 2009; Wilson and Sullivan 2011; Bolding and Franks 2017; Berners-Lee et al. 2023; Tantirigama, Huang, and Bekkers 2017). The percentage of cells in the food-specific subclass during slow feeding is comparable to odor-specific populations observed in the aPC (Tantirigama, Huang, and Bekkers 2017; Roland et al. 2017). When odors are presented at different intensities, the distributed population code and the neuronal firing rates remain similar (Bolding and Franks 2017). So, in spite of the higher food volume over time during binge feeding resulting in increased intensity of flavors of the same identity, we did not expect changes in the population code (Fig. 1g, 2E). Thus, our data points to a feeding rate-dependent modulation of flavor representation in the aPC that fundamentally alters the flavor-specific population code.

Flavor representation in the gustatory cortex is stable across feeding rates

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In general in sensory systems, behavior has profound effects on brain activity underlying the representation of sensory inputs (Musall et al. 2019; Steinmetz et al. 2019; Stringer et al. 2019). Therefore, neuronal suppression in the aPC induced by binge feeding could be a more global phenomenon that might be observed in other brain regions involving flavor representation. To test this, we performed Ca²⁺ imaging in the gustatory cortex (GC, granular, and dysgranular insular cortex) using a miniscope and tracked taste representation during slow and binge feeding 100 (Fig. 2A-C, S4A-G). We found little modulation in the general population of GC neurons upon 101 binge feeding compared with slow feeding (Fig. 2D), except for prolonged activation of GC neu-102 rons during binge feeding (Fig. 2D). in GC water-activated, non-selective and non-responding neurons, binge feeding of food had minimal levels of modulation compared with the general 104 suppression in aPC neurons. (Fig. 2E, Fig. S5A; estimated effect sizes in each neuron class for 105 aPC and GC neurons). Unlike the global activity suppression of the food-activated neurons in 106 the aPC, food-specific activation of GC neurons was selectively preserved during binge feeding (Fig. 2D, E, and S5B). 108

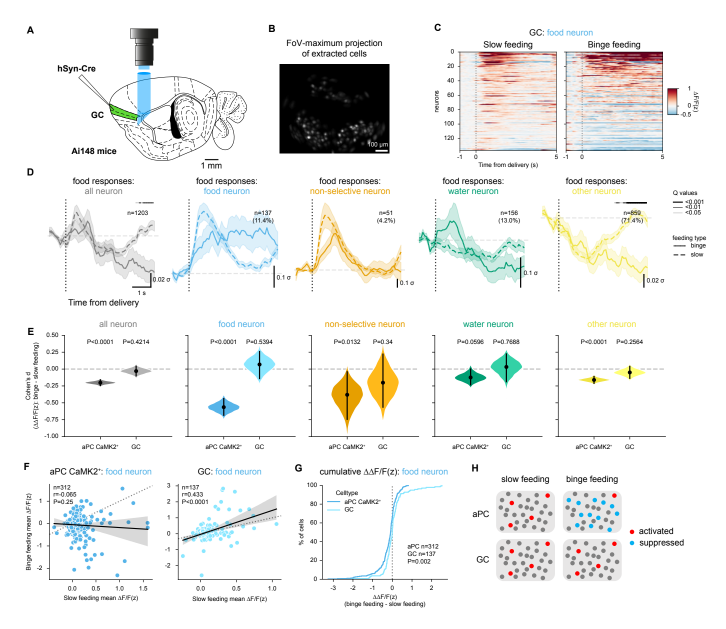


Figure. 2. Flavor representation in the GC is stable across feeding rates.

(A) Schematics of GRIN lens/Prism implantation in the GC. (B) Cell maps extracted by *CNMFe*. (C) Trial-averaged responses of individual GC food-activated neurons upon slow feeding and binge feeding (n=137 from 3 mice). (D) Trial-averaged responses of the whole population and subclasses of GC neurons upon slow and binge feeding (n=1203 cells from 3 mice). The shaded line above denotes the adjusted P-values (Q-values) of each time point, with different line widths representing different values (from thin to thick: Q <0.05, <0.01, <0.001). (E) Estimated effect size (Cohen's d) of binge feeding-induced modulation in the aPC CaMK2⁺ and GC neurons within individual subclasses. P-values are calculated with the permutation test with 5000 times bootstrapping (n is the same as in D). (F) Cell-wise comparison of neuronal responses upon slow feeding and binge feeding in food-activated aPC CaMK2⁺ and GC neurons (n is the same as in Fig. 1F and in C).

(G) Cumulative distribution of the difference (binge feeding vs. slow feeding) of z-scored $\Delta F/F$ ($\Delta\Delta F/F(z)$) in food-activated aPC CaMK2⁺ and GC neurons (n is the same as in **F**). (**H**) Schematics of neuronal responses in the aPC and the GC during slow feeding and binge feeding. For (**D**), data are shown as mean \pm s.e.m. For (**E**), data are shown as means of bootstrapped effect sizes (Cohen's d) \pm 95% confidence interval. For (**F**), r and P represent the correlation coefficient and P-value of Pearson's r. For (**G**), the P-value is calculated from the Kolmogorov-Smirnov test.

To further analyze these distinct modes of flavor representation across the two cortices, we correlated the single-cell activity between binge and slow feeding in food-activated neurons 111 in the aPC and the GC. In the GC, the activation level of food-activated neurons during slow feeding linearly correlates with the activity changes during binge feeding (Fig. 2F). In the aPC, 113 however, we found no correlation between responses during slow and binge feeding, suggesting that binge-induced activity modulation is independent of the neuronal responses during slow 115 feeding (Fig. 2F). The cumulative distribution of the amplitude difference between binge and slow feeding shows that in the GC, the net reduction is smaller and a larger fraction of cells show an increased binge feeding related neuronal response compared to the aPC (Fig. 2G). 118 We interpret our finding as a non-uniform, general inhibition during binge feeding (Frank et al. 119 2019). Thus, flavor representation in the GC is preserved during binge feeding, whereas binge 120 feeding induces a generalized activity suppression in the aPC (Fig. 2H).

Binge feeding-induced anterior piriform cortex suppression is not inherited from the olfactory bulb

The aPC is the first cortical relay of the olfactory system and receives sensory afferent inputs 124 from the olfactory bulb (Bekkers and Suzuki 2013). We tested if a reduction in sensory input 125 from the olfactory bulb could explain global suppression during binge feeding in the aPC. To 126 this end, we performed *in vivo* head-fixed 3-photon Ca²⁺ imaging in the olfactory bulb (OB) 127 mitral cells during binge feeding. Mitral cells are the major neuron population propagating odor 128 information to higher olfactory cortices, including aPC (Blazing and Franks 2020), and OB 129 mitral cells remained activated upon binge feeding (Fig. 3A-C, S6A, B). At the population level, net excitatory OB output to the aPC was similar during slow and binge feeding (Fig. 3D). These 131 findings align with the observation that intracortical connections and not the sensory afferents 132 dominate neuronal activity in the aPC (Poo and Isaacson 2011). We conclude that binge feedinginduced aPC suppression is not inherited from the OB.

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Binge feeding-induced anterior piriform cortex suppression extends to the major classes of local GABAergic neurons

Since the excitatory sensory drive is unaffected by feeding rate, enhanced recruitment of in-137 hibitory interneurons during binge feeding could underlie the global suppression of the aPC. 138 Local inhibitory feedback interneurons –activated by recurrent excitatory activity –predomi-139 nantly inhibit odor responses in the aPC, whereas the contribution of feedforward interneurons 140 is minor (Bolding and Franks 2018). We next probed the activity levels of aPC PV⁺ and SST⁺ 141 inhibitory interneurons during slow and binge feeding with miniscope Ca2+ imaging, as they 142 cover a large proportion of local feedback circuits (Large et al. 2016). Both PV⁺ and SST⁺ interneurons show strong suppression upon binge feeding. In contrast, the population responses 144 of both types of interneurons showed net activity increases during slow feeding (Fig. 3E-L, 145 S7A-F, S8A-C). Therefore, changes in the excitation-to-inhibition ratio of sensory afferent and 146 local recurrent aPC circuits do not seem to mediate the global suppression of aPC activity dur-147 ing binge feeding. The binge feeding-induced suppression of activity in the aPC affects both 148 the excitatory neurons and the major classes of local inhibitory interneurons. Therefore, binge 149 feeding-induced suppression globally affects most local circuits in the aPC. 150

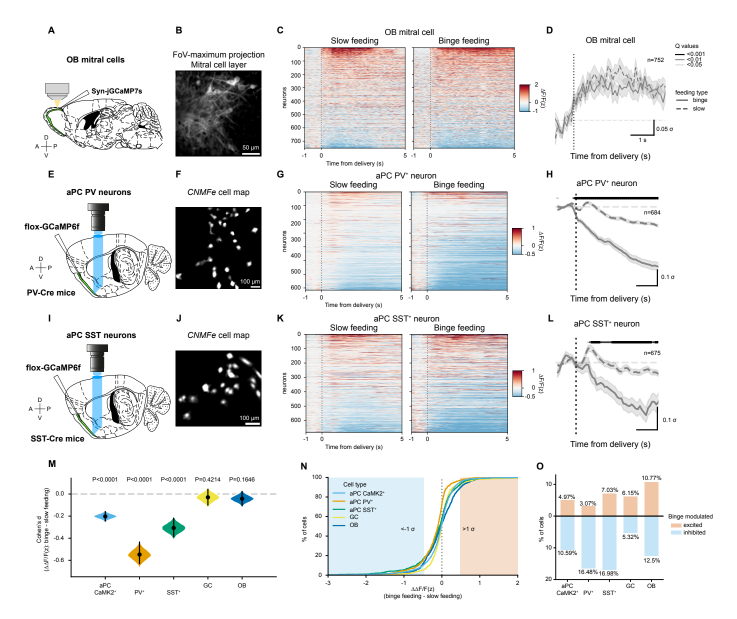


Figure. 3. Binge feeding-induced aPC suppression is not inherited from the OB and extends to GABAergic aPC neurons.

(A) Schematics of 3P-Ca²⁺ imaging in OB mitral cells. (B) FoV of OB mitral cells. (C) Trial-average of individual OB mitral cells upon slow and binge feeding (n=752 cells from 4 mice). (D) Trial-average of OB mitral cell population responses upon slow and binge feeding (n is the same as in C). (E) Schematics of Ca²⁺ imaging in aPC PV⁺ neurons. (F) Cell map of aPC PV⁺ neurons extracted by *CNMFe*. (G) Trial-average of individual aPC PV⁺ neurons upon slow and binge feeding (n=684 cells from 3 mice). (H) Trial-average of population aPC PV⁺ neuron responses upon slow and binge feeding (n is the same as in G). (I) Schematics of Ca²⁺ imaging in aPC SST⁺ neurons. (J) Cell map of aPC SST⁺ neurons extracted by *CNMFe*. (K) Trial-average of individual aPC SST⁺ neurons upon slow and binge feeding (n=675 cells from 3 mice).

(L) Trial-averaged of population aPC SST⁺ neuron responses upon slow and binge feeding (n is the same as in **K**). (**M**) Estimated effect sizes of binge-induced modulation in aPC CaMK2⁺, aPC PV⁺, aPC SST⁺, GC, and OB mitral cells. (**N**) Cumulative distribution of binge-induced modulation of $\Delta F/F(z)$ in aPC CaMK2⁺, aPC PV⁺, aPC SST⁺, GC, and OB mitral cells. (**O**) Percentages of binge feeding modulated population in aPC CaMK2⁺, aPC PV⁺, aPC SST⁺, GC, and OB mitral cells.

For (**D**), (**H**), and (**L**) data are shown as mean \pm s.e.m, and the shaded line above denotes the adjusted P-values (Q-values) of each time point, with different line widths representing different values (from thin to thick: Q <0.05, <0.01, <0.001). For (**M**), data are shown as means of bootstrapped effect sizes (Cohen's d) \pm 95% confidence interval.

Magnitude of binge feeding-induced anterior piriform cortex suppression correlates with appetite and depends on olfactory perception and metabolic state

Flavor perception of food items contributes to satiation and bypassing flavor perception via an 154 intragastric catheter reduces satiation and accelerates gastric emptying of identical food items 155 (Cecil, Francis, and Read 1999). The suppression of neuronal activity, specifically in the aPC 156 described here, reduces the sensory representation of food items. Accordingly, we hypothe-157 sized that the aPC suppression during binge feeding could constitute a mechanistic link between 158 a sensory neuronal response pattern in the flavor system and decreased satiation. Under ad li-159 bitum feeding conditions, mice consumed different amounts of food on different experimental 160 days, which we take as a proxy for differences in satiation. This noticeable behavioral variability in our recording sessions correlates with temporal progression, suggesting binge feeding 162 gradually escalates over time (Fig. S9A). We, therefore, investigated whether this behavioral variability maps onto the aPC neuronal responses. Using a linear mixed model, we found a 164 robust time-independent correlation between the initial binge eating-induced aPC suppression and subsequent food consumption on each recording session (Fig. 4B). Suppression was al-166 ways quantified during the onset of a binge bout within the first 4 seconds after initiation. Our 167 model, therefore, quantifies suppression independently of feeding duration. The strong corre-168 lation between neuronal activity patterns and food intake does not exist for slow feeding aPC 169 responses (Fig. 4B, C). We also observed consumption-correlated suppression of neuronal ac-170 tivity in aPC PV+ neurons. In contrast, aPC SST+ neurons and GC neurons did not show a 171 correlation between neuronal activity and food intake. (Fig. S9C). These findings suggest that 172 consumption-correlated neuronal modulations are mostly restricted to the olfaction component of flavor perception.

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We further examined the necessary factors for the consumption-correlated generalized suppression of aPC neuronal activity during binge feeding. To test whether the olfactory perception of 177 the food items is a necessary for consumption-correlated suppression of neuronal activity, we 178 performed nasal lavage with 0.5% Triton solution to induce temporary anosmia in mice (Cum-179 mings et al. 2000) (Fig. 4D, E, anosmia verified by buried food test). We performed imaging 180 experiments before and after the intervention. Our procedure caused a 30% drop on average 181 in the number of active neurons in the field of view detected with imaging, consistent with re-182 duced olfactory inputs to the aPC (Fig. S9B). We then compared the correlation of the aPC 183 activity suppression to consumption in the same mice before and after the anosmia-inducing 184 treatment. In contrast to the pre-anosmia condition, anosmia-inducing treatment abolishes the 185 correlation between the suppression of aPC neuronal activity and consumption (Fig. 4 F, G). Un-186 der anosmia, we still observed binge eating-induced aPC suppression (Fig. 4H) in the presence 187 of generally enhanced food intake (Fig. 4F). This result suggests that intact sensory olfactory 188 perception is a prerequisite for binge eating-induced aPC suppression correlated to consump-189 tion. Metabolic states like hunger and satiety profoundly affect sensory systems (Soria-Gómez 190 et al. 2014; Aimé et al. 2007; Freeman 1960; Prud'homme et al. 2009; Albrecht et al. 2009). 191 Consequently, we wondered if changes in the metabolic state affect the consumption-correlated 192 suppression of aPC neuronal activity during binge feeding. We altered the metabolic state of 193 mice by overnight fasting. We found enhanced food intake in these mice and enhanced binge feeding-induced suppression of aPC neuronal activity compared to ad libitum-fed conditions 195 (Fig. 4I-L). Under fasting, the correlation between food consumption and aPC neuronal activity suppression is lost (Fig. 4J). An increase in suppression was also observed in the GC of fasted 197 mice (Fig. 4K, L). We conclude that the suppression of neuronal activity during binge feeding in the aPC is enhanced by fasting and is also observed in the GC.

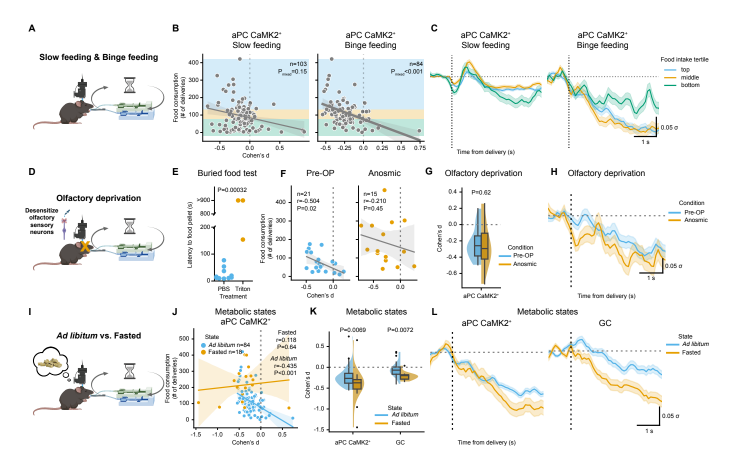


Figure. 4. Magnitude of binge feeding-induced anterior piriform cortex suppression correlates with appetite and depends on olfactory perception and metabolic state.

(A) Schematics of feeding and Ca²⁺ imaging paradigm. (B) Correlations between session-specific food intake and modulation of neuronal activity in aPC CaMK2⁺ neurons upon slow and binge feeding (n=103 slow feeding sessions and 84 binge feeding sessions from 8 mice). (C) Trial-average responses of aPC CaMK2⁺ neurons upon slow and binge feeding clustered by food intake (n=34, 35, 34 sessions for top, middle, bottom clusters in slow feeding from 8 mice, n=28, 28, 28 sessions for top, middle, bottom clusters in binge feeding from 8 mice). (D) Schematics of anosmic paradigm. (E) Latency of mice finding the buried food pellet 48 hrs after treatment (n=10 mice for PBS treated group, n=3 for Triton treated group). (F) Correlations between session-specific food intake and modulation of neuronal activity in aPC CaMK2⁺ neurons upon binge feeding with intact olfaction and under anosmic conditions (n=21 Pre-OP sessions and n=15 Anosmic sessions from the same 3 mice). (G) Binge-induced modulation of neuronal activity with intact olfaction and under anosmic conditions (n is the same as in F). (H) Trial-averaged activity of aPC CaMK2⁺ neurons upon binge feeding with intact olfaction and under anosmic conditions (n is the same as in F). (I) Schematics of fasting paradigm. (J) Correlations between session-specific food intake and modulation of neuronal activity in aPC CaMK2⁺ neurons upon binge feeding under ad libitum or overnight fasted conditions (n=18 fasted sessions from 5 mice, n for ad libitum conditions is the same as in B). (K) Binge-induced modulation of neuronal activity under ad libitum or overnight fasted conditions (n is the same as in **J**).

(L) Trial-averaged responses of aPC CaMK2⁺ neurons upon binge feeding under *ad libitum* or overnight fasted conditions (n is the same as in **J**).

For (C), (H), and (L) data are shown as mean \pm s.e.m. For (F) and (J) r and P represent the correlation coefficient and P-value of Pearson's r. For (B) P represents the significance level of Cohen's d to food consumption by a linear mixed model. For the box plot in G and (K) the center line shows the median, the box limits show the quartiles, the whiskers show 1.5x the interquartile range, and the points show the outliers.

Optogenetically suppressing anterior piriform cortex neurons promotes feeding

We have so far established the correlative relationship between feeding rate, olfactory flavor rep-201 resentation, and metabolic state. While binge feeding and appetite clearly covary with global 202 activity suppression in the aPC, it is unclear if the sensory effect we observe is an epiphenomenon 203 (H0, Fig. 5A) or reciprocally interacts with feeding behavior in a feedback loop (H1, Fig. 5A). 204 We, therefore, next asked whether there was a causal relationship between suppressed aPC activity and feeding behavior by inhibiting aPC during feeding. This tested if aPC suppression 206 alone is sufficient to increase food consumption (H1, Fig. 5A). We employed a closed-loop op-207 togenetic inhibition paradigm to silence aPC excitatory neurons at the initiation of binge feeding 208 bouts (Fig. 5B, E). To suppress activity at the behavioral timescale (tens of seconds) of binge 209 feeding bouts while minimizing the illumination period, we chose the highly light-sensitive 210 mosquito opsin eOPN3 (Mahn et al. 2021) to provide long-lasting suppression of recurrent ex-211 citatory fibers in the aPC (Fig. S10C for optical fiber implant coordination). We found that mice 212 consumed more food when aPC activity was optogenetically suppressed upon feeding (Fig. 5F-213 I). The optogenetic suppression of aPC activity prolonged the individual feeding bouts, while 214 the number of feeding bouts remained similar, suggesting that optogenetic aPC suppression 215 predominantly affects consummatory and not appetitive behavior (Fig. 5F and Fig. S10A, B). Light stimulation alone did not affect feeding behaviors in control mice transduced with AAVs 217 encoding tdTomato (Fig. 5G-I). Our data infers that binge feeding-induced aPC suppression is 218 causally linked to feeding behaviors, suggesting a functional role of binge feeding-induced aPC 210 suppression in modulating appetite (*H1*, Fig. 5A)

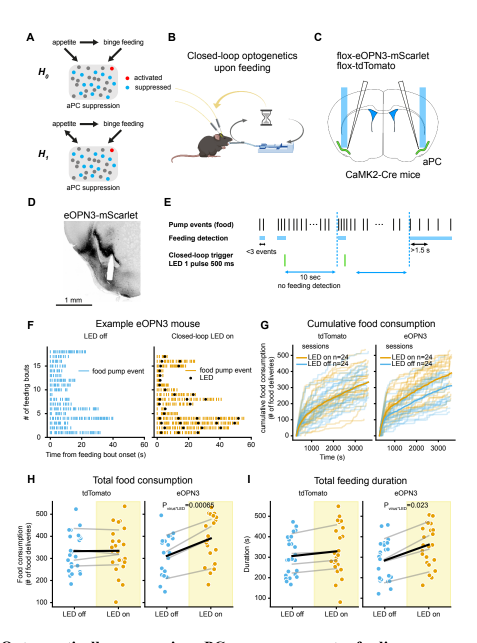


Figure. 5. Optogenetically suppressing aPC neurons promotes feeding.

(A) Schematics of hypotheses on relationships of appetite, binge feeding and aPC suppression. (B) Schematics of closed-loop optogenetics experiment setup and feeding paradigm. (C) Schematics of viral injection and optical fiber implants bilaterally in the aPC. (D) Brain slice with optical fiber implant path. (E) Schematics of feeding-based closed-loop optogenetics paradigm. (F) Example feeding bouts in an eOPN3-expressing mouse without light stimulation (left panel) and with closed-loop light stimulation (right panel). (G) Cumulative feeding events through experimental sessions for mice expressing tdTomato (left panel, n=12 LED off and 12 LED on sessions from 4 mice) and mice expressing eOPN3 (right panel, n=12 LED off and 12 LED on sessions from 4 mice). (H) Effects of light stimulation on total food consumption in tdTomato- and eOPN3-expressing mice (n is the same as in G).

(I) Effects of light stimulation on total feeding duration in tdTomato- and eOPN3-expressing mice (n is the same as in G). For (H) (I) grey lines denote the data from the same mice, and the black line denotes the mean. The P-values are calculated from a linear mixed model (see **Methods**).

DISCUSSION

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The role of olfactory flavor representation during feeding behaviors is poorly understood despite its essential contribution to flavor experience (Shepherd 2006; Maier et al. 2015). Using cell-type specific *in vivo* Ca²⁺ imaging and optogenetics in freely behaving mice, we provide circuit-level evidence that suppression of flavor representation in the aPC during binge feeding actively enhances food intake. While chronic effects of olfactory alterations on food intake and metabolism have been reported (Riera et al. 2017; Tucker, Overton, and Fadool 2012), we found an acute functional role of olfaction at the level of individual feeding bouts. Our findings suggest that the olfactory representation of flavor during feeding provides feedback for sensory satiation to modulate food intake and homeostasis in real-time.

Metabolic states have profound effects on sensory systems, especially olfaction. Hunger increases neuronal and behavioral responses to odors, and in contrast, satiety decreases them (Soria-Gómez et al. 2014; Aimé et al. 2007; Freeman 1960; Prud'homme et al. 2009; Albrecht et al. 2009). Sensory detection of food items leads to a rapid switch in activity patterns from consumption-promoting AgRP neurons to consumption-inhibiting POMC neurons in the arcuate nucleus of the hypothalamus. The amplitudes of these foraging-related switches in hypothalamic activity are enhanced by fasting and sensory signals from food items with a high hedonic value (Chen et al. 2015; Betley et al. 2015; Mandelblat-Cerf et al. 2015). We demonstrated that the amplitude of binge feeding-induced suppression in the aPC reflects internal appetite levels and is enhanced by fasting, supporting that metabolic states strongly modulate olfactory representation during feeding.

The sensory experience of food items contributes to satiety. Bypassing sensory experiences of food within the oral cavity by direct gastric infusion reduces satiety and accelerates gastric emptying compared to regular feeding in humans and rodents (Berkun, Kessen, and Miller

²⁴⁶ 1952; Cecil, Francis, and Read 1999; Stratton and Elia 1999). A recent study also demonstrates a brainstem circuit of gustatory oral sensory feedback that induces satiation (Ly et al. 2023). Our study supports the idea that suppressed olfactory flavor representation, via direct binge feeding or artificially suppressing the aPC, constitutes an acute intrinsic behavior-associated cortical mechanism leading to more consumption, inferring lower satiety levels. The reciprocity between feeding behavior and olfactory representation demonstrated here (Fig. 5a) extends the emerging theory that perception and action reciprocally interact (Buzsáki 2019).

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From an evolutionary perspective of food scarcity, overeating induced by suppression of sensory satiety would be pro-survival. In the surplus of food environment human beings are facing today, an increased eating rate is commonly linked to overeating (Hall et al. 2019) and obesity (Ohkuma et al. 2015), while reducing the eating rate can effectively mitigate food consumption (Hurst and Fukuda 2018; Scisco et al. 2011; Bolhuis et al. 2014). Here, we provide evidence that feeding rate-dependent modulation of olfactory flavor representation modulates appetite (Fig. S11). Our findings add the perspective of sensory experience and awareness to the notion that it is not only what you eat but also how you eat.

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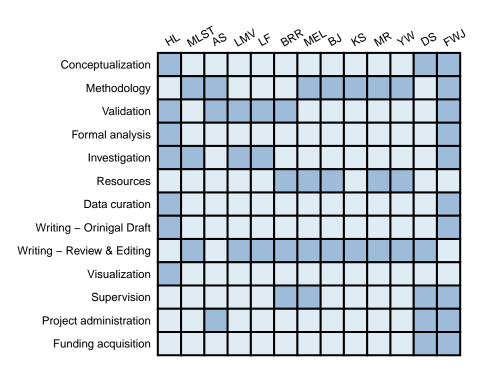
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ACKNOWLEDGEMENTS:

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We thank Susanne Rieckmann, Monika Dopatka, Anny Kretschmer, Katja Czieselsky and Celina
Ernst for their excellent technical assistance; Daniel Parthier for his inputs on statistical analysis;
Yen-Chung Chen for his inputs on programming; Melissa Long from the Charité Animal Behavioral Phenotyping Facility, Charité Advanced Medical BIOimaging Core Facility, and Charité
Viral Core Facility for their assistance; James Poulet and David Owald for their feedback on
the manuscript; Robin Blazing and Kevin Franks for sharing *in vivo* optical fiber implantation
protocols; Andreas Klaus for his input at the early stage of this study.

HL is supported by the PhD Fellowship from the Einstein Center for Neurosciences Berlin and 438 the Taiwanese Government Scholarship to Study Abroad. FWJ is supported by the Deutsche 439 Forschungsgemeinschaft (DFG) (project 458236353 and project 520223756 - SPP2411 LOOPs). 440 BRR is supported by the DFG (project 327654276 - SFB1315 and project 273915538 - SPP1926). 441 DS is supported by the DFG (project 184695641 –SFB 958, project 327654276 –SFB 1315, 442 project 415914819 -FOR 3004, project 431572356, the German Excellence Strategy EXC-443 2049-390688087, NeuroCure), by the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation program (BrainPlay Grant agreement No. 810580) 445 and by the Federal Ministry of Education and Research (BMBF, SmartAge – project 01GQ1420B).



Author contribution

AUTHOR CONTRIBUTIONS

Conceptualization: HL, DS, and FWJ; methodology: AS, FWJ, KS, MR, YW, MLST, BJ, and MEL; validation: AS, HL, LF, LMV, BRR, and FWJ: formal analysis: HL and FWJ; investigation: HL, FWJ, MLST, LF, and LMV; resources: MEL, BJ, MR, YW, DS, and BRR; data curation: HL and FWJ; writing - original draft: HL and FWJ; writing - review & editing: DS, BRR, MR, BJ, KS, MLST, MEL, LF, LMV, and YW; visualization: HL; supervision: FWJ, DS, MEL, and BRR; project administration: AS, FWJ, and DS; funding acquisition: FWJ and DS.

DECLARATION OF INTERESTS

454 YW owns PhenoSys equity. Other authors have declared no competing interest.

DATA AND CODE AVAILABILITY

- Source data for individual figure panel can be found on a GitHub repository
- (https://github.com/hung-lo/BingeFeeding_2023). Raw data will be available upon request due
- to the large file size (~2-3 TB). Code for plotting individual figure panel can be found on a
- GitHub repository (https://github.com/hung-lo/BingeFeeding 2023). All code for data process-
- ing and analysis will be deposited to a GitHub repository upon acceptance of this manuscript.

METHODS

Animals

Animals were kept at the animal facility of Charité, under a regular 12/12 hour light-dark cycle. 461 All procedures involving animal experiments were approved by the local authorities and ethics 462 committee (LaGeSo Berlin, license numbers G0278/16, G0313/16, and G0156/20). To image 463 excitatory neurons in the aPC, we cross-bred Ai93D mice, Rosa-tTA mice, and CaMK2-Cre mice to obtain Ai93D; Rosa-tTA; CaMK2-Cre mice that express GCaMP6f in excitatory cells. 465 To prevent early expression of GCaMP during development, the breeding pairs and offspring 466 were fed with Doxycycline-containing food to suppress the expression of GCaMP6f until wean-467 ing. Due to suboptimal GCaMP6f expression in the GC in the abovementioned transgenic mice, 468 we injected the AAV virus carrying hSyn-Cre in the GC of Ai148D mice to express GCaMP6f 469 in the GC. To image OB mitral cells, we inject Syn-jGCaMP7s AAV virus in C57BL/6 mice. To 470 image GABAergic neurons in the aPC, we performed viral injection of Cre-dependent GCaMP6f 471 in PV-Cre or SST-Cre mice. For optogenetic experiments, CaMK2-Cre mice were bilaterally 472 injected with Cre-dependent eOPN3 virus or a Cre-dependent tdTomato expressing virus for controls. All experiments including Ca²⁺ imaging and optogenetics are performed between 9 474 a.m. to 6 p.m. under regular light.

Liquid food delivery system

To reduce stress, we performed experiments inside the animals' home cages. Cages were mod-477 ified so that we could protrude the motorized lick spout (PhenoSys, Berlin, Germany) into the 478 cage. After a 5-minute baseline period, motorized lick spouts were presented in the cage and 479 primed for liquid delivery for 2 min with 1 min intervals between presentations. During these intervals, the spouts were retracted. For olfactory isolation, we presented the lick spout inside 481 a glass tube with an opening for animals to reach the spout. Air suction from the glass tube 482 limited olfactory responses to the food odor to the time period just before and while mice inter-483 acted with the lick spout. Lick spouts were equipped with piezo sensors to register each licking. 484 Licks triggered a 400-ms activation of electrical pumps, which resulted in the delivery of one 485 droplet (~1.8 μL) of strawberry or chocolate-flavored Ensure (Abbott Laboratories) or water. After delivery, we set a refractory delay period of 4 sec (slow feeding) or 400 ms (binge feeding). During slow feeding, we provided Ensure and water at a ratio of 7 to 3. Four slow-feeding rounds and three binge-feeding rounds were interchanged in a pseudorandom order. Mice were *ad libitum* fed before the experiments, with a maximal period of up to 4 hours of pre-experiment food deprivation during the light cycle. Fasting was performed once a week for 20-22 hours before starting the experiment.

Surgery procedures

94 Stereotactic injection

Mice were anesthetized by inhalation anesthesia with isoflurane (induction: 4-5%, then 1-2% 495 with oxygen, flow rate 0.5-1 l/min). Mice were local anesthetized with Lidocaine (1-2%) subcutaneous injection preincision. A craniotomy was performed over the stereotactically determined 497 target regions (Table. S1) using a semi-automatic neurostar stereotactic apparatus (Neurostar, Tübingen, Germany). The virus (0.4 to 1 μL) was injected using a 10 μL-Hamilton syringe. 490 Postoperative pain was prevented by Carprofen (5 mg/kg) subcutaneous injection right before 500 surgery and in the first 3 days after surgery. After the surgery, the animals recovered for at least 501 two weeks. In some experiments, implantation of the prism or the optic stimulation fiber was 502 performed right after viral injection. 503

504 GRIN lens implantation

Mice (>P50) were anesthetized for the procedure by inhalation anesthesia with isoflurane (induction: 4-5%, then 1-2% with oxygen, flow rate 0.5-1 L/min). Mice were local anesthetized 506 with Lidocaine (1-2%) subcutaneous injection preincision. The anesthetized animals were fixed 507 in the stereotact (Neurostar, Tübingen), and a craniotomy was performed over the stereotacti-508 cally determined target region (Table. S1). The side length of the quadratic craniotomy was 509 slightly larger than the side length of the prism base, approximately 1.2 mm. The insertion tract 510 was paved by aspiration of brain tissue until ~ 1 mm above the image plane of the microscope. 511 Aspiration was performed through a thin needle (23G, sharp end) linked to a vacuum pump, the 512 procedure was performed twice to ensure sufficient aspiration of brain tissues. Any small hem-513 orrhagic foci that occurred were staunched by Gelfoam. After removal of the Gelfoam and any

pending blood clots, insertion of the microendoscopic lens (GRIN lens attached with a Prism (1 mm diameter, \sim 9.1 or \sim 4.3 mm long, Inscopix) to the desired image plane is performed at 516 a rate of 100 μm/min according to the coordinates in Table S1. With the aid of an adhesive (VetBond, 3M or TRUGLUE, TRUSETAL) and dental cement (Super-Bond C&B, SUNMED-518 ICAL), microendoscopes with attached baseplates (Inscopix) were fixed to the skull, and the 519 optical surface was protected from contamination by a plastic cap (Inscopix baseplate cover). 520 Postoperative pain was prevented by Carprofen (5 mg/kg) subcutaneous injection right before 521 surgery and in the first 3 days after surgery. In a subset of mice, we did not use microendoscopes 522 with attached baseplates. In this case, the slightly protruding microendoscope was fixed to the 523 skull with adhesive and dental cement, and the optical surface was protected from contamina-524 tion using a silicone cap. In these mice, a baseplate was fixed in the desired optical plane above 525 the protruding lens in a surgery that was performed at a minimum of four weeks after the lens implant. 527

528 Cranial window for olfactory bulb imaging

Mice (C57BL/6, <P40) were anesthetized with isoflurane (induction: 4-5%, then 1-2% with 529 oxygen, flow rate 0.5-1 l/min). Mice were local anesthetized with Lidocaine (1-2%) subcuta-530 neous injection preincision. After the scalp and periosteum were removed, a 3mm craniotomy was made over the two bulb hemispheres. An injection micropipette (tip diameter, $10-20 \mu m$) 532 was filled with AAV1.Syn.jGCaMP7s.WPRE virus solution (Penn Vector Core), and 100 nl was injected 50 nl/min at a depth of 300 μ m in either bulb hemisphere (see Table S1 for coordinates). 534 After injection, a semi-circular <3 mm stack of two glass coverslips, glued to each other using 535 optical adhesive, was fitted into the craniotomy and sealed with cyanoacrylate glue and dental 536 cement. Finally, a light-weight head-post was fixed on the skull over the left hemisphere with 537 light-curing adhesives (RelyX, 3M) and dental cement (Ortho-Jet, Lang Dental). Postoperative 538 pain was prevented by Buprenorphine (0.05-0.1 mg/kg) and Carprofen (5 mg/kg) subcutaneous 539 injection right before surgery and then Carprofen (5 mg/kg) in the first days after surgery. Headfixed 3-photon imaging experiments began 3 weeks after the virus injection.

Ca²⁺ imaging

Habituation

550

For freely-moving recordings, before starting the combined behavioral and imaging sessions using Ensure and water, mice were habituated to the lick spout delivery with 10% sucrose solution. Mice had to reach a criterion of 25 sucrose deliveries in slow feeding mode in a 45-minute habituation session before the actual measurements began. Mice were further habituated with additional air suction around the lick spout and dummy scope mounting once they had learned to drink from the lick spout. The habituation period usually lasted ~3 weeks.

For head-fixed recordings, the habituation of mice to head-fixation began at least 5 days prior to imaging. On the first day, the animal was head-fixed on a running wheel for 5 min and then gradually increased each day until it was calm for 1 hour. At least one day before imaging, a lick spout with milk/water within easy reach for licking was introduced.

555 in vivo imaging: miniscope

The miniaturized microscope (nVista miniscope, Inscopix, CA, USA) was mounted right before 556 the imaging session started without anesthesia. Before the recording started, mice were allowed 557 to explore the home cage for 3-5 minutes. After a baseline period of 5 min, the lick spout 558 protruded according to the protocol described above. For each mouse, the imaging settings 559 (LED intensity, gain, focus ··· etc.) were individually tuned to reach a similar level of brightness 560 (mean values around 50-60 A.U. in fluorescence histogram function in Inscopix acquisition 561 software). We recorded at 20 Hz with a single focal plane. Most imaging sessions were 4x 562 spatially down-sampled during acquisition to save storage space. The behavioral system was 563 linked with the Inscopix system using TTL pulses upon pump activation. We performed up to 25 imaging sessions per mouse across 5 weeks, and overnight food deprivation was performed once per week.

in vivo imaging: 3-Photon

Imaging from head-fixed mice was performed with a home-built 3-photon microscope. The laser 568 (Opera-F, pumped by Monaco, Coherent) provided light pulses at 1300 nm wavelength and 1 MHz repetition rate for excitation of jGCaMP7s. The laser output passed a four-pass prism 570 pulse compressor for dispersion compensation. Laser power was adjusted using a motorized 571 half-wave plate and a polarizing beam splitter, and was below 20 mW under the objective. We 572 used a Nikon25x/1.1 objective and dual linear galvanometers at a frame rate of \sim 10Hz. Image 573 acquisition was synchronized with laser pulses and was controlled by LSMAQ. Time-series 574 images (200 x 200 pixels) were recorded at depths of 200-300 μ m below pia at the mitral cell 575 laver. 576

577 Triton X-100 application

Nasal lavage with 0.5% of Triton X-100 (in 0.1M PBS) can introduce temporal anosmia in mice for up to 3 weeks (Cummings et al. 2000). Mice were anesthetized with Ketamin (100 579 mg/Kg), Xylazine (20 mg/Kg) and Acepromazine (3 mg/Kg) intraperitoneal injection. They 580 received Caprofen (5mg/Kg) subcutaneous injection before and the day after the Triton X-100 581 (experimental group) or PBS (control group) applications. We applied 40 μL 0.5% Triton X-582 100 solution to each nostril with a gel loading pipette tip that was advanced for 2-3mm into the 583 nostril. Triton solution was slowly applied with a micropump (Narishige, Japan) over several 584 minutes on each side, with an interval of 5 min between the two nostrils. Foam building up at 585 the opening of the nostrils was an indicator of a successful procedure. Throughout the procedure 586 and until waking up from anesthesia, mice were kept on an inclined plane so that their nostrils were below their trachea and lungs in a heated chamber. 588

Buried food test

To test the efficacy of Triton-induced anosmia, we examined mice's ability to find a hidden food pellet located 1 cm deep in the bedding of the experimental cage. Mice were overnight food-deprived and habituated to the experimental cage for at least 5 minutes before the experiment started. After the food pellet was buried, mice were transferred into the experimental cage. The time to find the pellet was documented by the experimenter. If the mice did not find the pellet

after 15 minutes, the experiment was stopped. Control experiments were performed with mice that had undergone the same lavage procedure with 0.1M PBS nasal lavage. The buried food tests were repeated every week to monitor mice's smell ability to ensure mice remained anosmic throughout our experiments.

Optogenetic experiments

Mice expressing eOPN3 in aPC excitatory neurons were anesthetized for the procedure by inhalation anesthesia with isoflurane (induction: 4-5%, then 1-2% with oxygen, flow rate 0.5-1 601 L/min). The anesthetized animals were fixed in the stereotact (Neurostar, Tübingen), and a craniotomy was performed over the stereotactically determined target region. Fiberoptic cannulas 603 (200 µm diameter, NA 0.66, length 5mm, Doric lenses) were inserted bilaterally until reach-604 ing the target coordinates (Fig. S10c). The slightly protruding fiber with an attached zirconia 605 sleeve for taking up the stimulation fiber patchcord was fixed to the skull with adhesive and 606 dental cement, and the optical surface was protected from contamination using a plastic cap. 607 Before starting the combined behavioral and optogenetic stimulation sessions using Ensure, the 608 mice were habituated to the lick spout delivery with 10% sucrose solution. Mice had to reach a 609 criterion of 25 sucrose deliveries in binge feeding mode and needed to successfully trigger sham 610 closed-loop stimulations in a 20-minute habituation session before the actual measurements began. At the start of the experiment, we plugged a splitter branching patchcord (200 µm diameter, 612 NA 0.57, Doric) connected to a mono fiberoptic patchcord (480 µm diameter, NA 0.63, Doric) onto the fiberoptic cannulas. Light from a Ce:YAG fiber light source (Doric) was delivered at 614 an intensity of 8mW at the tip of the fibers. For these experiments, mice were granted constant 615 access to the lick spout in the binge feeding mode over a period of 60 min. Upon detection of 616 a binge bout (3 pump deliveries with a maximum of a 1.5 s interdelivery interval), a 500 ms 617 light stimulus was delivered on "LED on" days. After the detection of a binge bout with a 618 subsequent light stimulus, there was a minimal refractory period of 10 seconds until a binge 619 bout could initiate the next light stimulus. Everything was similar on "LED off" days, apart 620 from not stimulating with light upon binge bout detection. LED on and off days were alternated 621 for 12 subsequent days.

Imaging processing

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Ca²⁺ movies obtained from miniscope recordings were first temporally downsampled to 10 Hz. We then cropped out regions in the field of view (FoV) where no active Ca²⁺ transients were visible. The same FoV cropping parameters were used throughout recordings from the same mice. 626 Movies were then bandpassed with a spatial filter (low cutoff=0.005, high cutoff=0.500, Inscopix IDPS) and motion corrected (aligned to mean image or first frame, max translation=20, 628 Inscopix IDPS). Ca²⁺ traces were extracted with an adapted version of *CNMFe* (Zhou et al. 620 2018) from Inscopix (see Table S4) with the following parameters (Cell diameter: 10 px, PNR: 630 10 for excitatory cells and 20 for GABAergic cells, Corr. 0.8). Ca²⁺ traces were manually cu-631 rated with predefined selection criteria (peak amplitude >80 A.U., baseline drifts less than 20% 632 of peak fluorescence, clear cell shape, locate outside blood vessels, minimal motion artifacts of 633 given regions of interest).

Image stacks from 3P imaging were loaded into Suite2P (Pachitariu et al. 2016) for motion correction, region-of-interest (ROI) segmentation and trace extraction using the settings specified in the appended exemplary .ops file. We used the 'mean img', 'correlation map', and 'max projection' views of Suite2P to manually check and sort somatic from non-somatic ROIs of mitral cells. The output from Suite2P was analyzed in Python: Detected neuropil signals were subtracted. Remaining frames with movement artifacts were then detected and excluded based on the presence of post-registration x- and y-shifts at each time point. A further criterion was the phase correlation of individual frames and the reference image below a threshold of 50% of the maximum peak of phase correlation in the respective stack. After that, $\Delta F/F$ values and z-scores were calculated in customized Python scripts.

Data analysis, statistics and plotting

All data analysis and statistics were performed using custom scripts in Python and R. Most figures were plotted in Python (matplotlib, seaborn) and figure and font sizes were later modified in Illustrator (Adobe). We used the Okabe-Ito color palette (Ichihara et al. 2008) to increase the accessibility of common forms of color blindness.

Data synchronization

Ca²⁺ imaging data and behavioral data were synchronized by finding the time lag of maximum cross-correlation between pump events in digital values from the PhenoSys behavioral protocol and the binarized pump-triggered TTL pulses recorded in the Inscopix system by Pynapple.cross_correlogram (Python).

Binge feeding bout detection and slow feeding processing

To detect binge feeding bouts, inter-pump intervals were calculated for each pump event, and only pump deliveries with intervals shorter than 2 s qualified as part of a feeding bout. Additionally, each feeding bout was required to include at least 3 pump deliveries.

Pump events in slow feeding mode were filtered out if no further lick event followed the initial lick event triggering pump activation. Initial motor artifacts from the movement of the lick spout were also removed from the further analysis. Since binge feeding bouts were guaranteed to have subsequent lick events by design, the exclusion of pump events following no lick events was not applied to binge feeding pump events.

665 Statistical analysis

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Statistical analysis was performed in Python (Scipy, Numpy, Dabest) and R (Lmer). Individual statistical tests are listed under the respective figure legends and all statistical details are listed in the statistical summary Table S5. All statistical tests are performed with two-sided testing. Mean \pm standard error of the mean (s.e.m.) or 95% confidence of interval were used to report statistics in figures. Applied statistical tests and the sample size for each analysis are listed in the figure legends and the statistical summary table. A significance level of p <0.05 is used for rejecting null hypothesis testing. No statistical methods were used to predetermine sample size randomization nor blinding was applied.

674 Area Under the Receiver Operating Characteristics (auROC)

We used auROC to classify neurons into different response classes to Ensure and water deliveries during slow feeding (Botta et al. 2020; Cohen et al. 2012). In individual neurons, we

compared the distribution of raw Ca²⁺ amplitudes during baseline activity (-1 to 0 s before pump activation) to the distribution of raw Ca²⁺ amplitudes in individual 100-ms bins across trials. To 678 produce the bin-specific ROC curves, we moved a criterion from the minimal to the maximal Ca²⁺ response we found in the neuron's baseline activity distribution and the given 100 ms 680 bin distribution. We then plotted the probability that Ca²⁺ signals in the given 100ms bin distri-681 bution were larger than the criterion against the probability that the Ca²⁺ signals in the baseline 682 distribution were larger than the criterion. The auROC for each bin was then calculated using 683 the auc function (sklearn.metrics.auc), resulting in auROC values between 0-1 and 0.5 means 684 not different from the baseline. The post-stimulus auROC values from each time bin were com-685 pared to the baseline auROC values. Significance was established if at least four consecutive 686 post-stimulus bin values between 0-2 s were greater than 2 S.D. of the pre-stimulus baseline 687 values (food- or water-activated neurons).

689 Effect size calculation

To calculate the effect size of binge feeding-induced suppression in each neuron class, we performed a bootstrap-coupled estimation (DABEST, Python). To obtain the distribution of the mean difference between the two conditions, we re-sampled the mean Ca²⁺ activity of two conditions (eg. slow feeding and binge feeding) 5000 times (bootstrapping distribution, represented as the violin plot in Fig. 2e, 3m, and Fig S8a). The distribution was then normalized by the pooled standard derivation of both conditions to convert it to Cohen's d using the Dabest.cohens_d (DABEST). P values were computed with the Dabest.PermutationTest (DABEST).

of O values calculation

To estimate the differences between 2 neuronal time series along the time axis, we first calculated the P values of each time bin by performing the unpaired Student's t-test (scipy.stats.ttest_ind) and then applied false discovery rate correction (statsmodels.stats.multitest.fdrcorrection) to obtain the adjusted P values, which are the Q values.

Linear Mixed Models

Linear mixed models were used to estimate contributions of predictors (eg. level of excitation/suppression in neuronal activities upon feeding, or interaction of optogenetic actuators and
light stimulation) to outcome (eg. food consumption or feeding bout duration) while allowing
different intercepts for individual mice (lmer, R). To calculate the contribution of a given predictor, we built a full model with all predictors and a reduced model that lacks the given predictor.
We then compared these two models with anova function (R) to calculate the P value of the
given predictor. Representative models are structured as following:

711 Full model

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$$Food.consumption \sim 1 + session.number + bodyweight + sex + \\virus.type + LED.state + virus.type * LED.state + (1|mouse.id)$$

714 Reduced model

$$Food.consumption \sim 1 + session.number + bodyweight + sex + \\ virus.type + LED.state + (1|mouse.id)$$

Histology and imaging

Mice were anesthetized with (100 mg/Kg) Ketamine and (15 mg/Kg) Xylazine and perfused with 0.1M PBS and then 4% paraformaldehyde (PFA). Mice brains were harvested and stored in 4% PFA at 4 °C overnight and transferred to 0.1M PBS for long-term storage. Brains were embedded in 4% agar-agar and sliced at 100-150 μm thickness with a vibratome. Brain slices were mounted on glass slides and were imaged by an epifluorescence microscope (Leica DMi8) or a confocal microscope (Leica SP5). Acquired images were then aligned to the mouse brain atlas (Paxinos and Keith B. J. Franklin 2007) for registration of the location of GRIN lens-prism or fiber optic cannula implants.

Serial 2-Photon tomography (Brainsaw)

A subset of fixed mice brains was sliced and imaged by serial 2P tomography, where whole forebrain structures can be imaged at cellular resolutions (Ragan et al. 2012). We modified a custom-made 2P microscope (COSYS, UK) to operate with BakingTray (ScanImage & BakingTray, MATLAB). Obtained images were stitched (StitchIt, MATLAB) and reconstructed into 3D brain models. Image stacks were then registered to the Allen mouse brain atlas (BrainReg, Python) and visualized with napari (Python).

SUPPLEMENTARY FIGURES

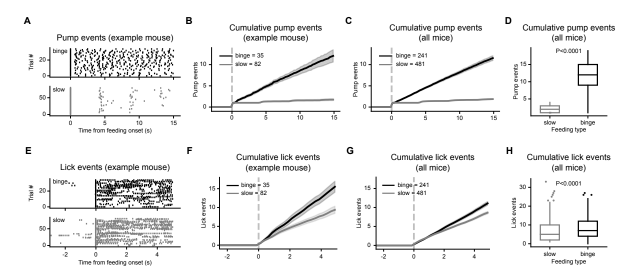


Figure S1. Behavioral differences between slow feeding and binge feeding.

(A) Pump delivery events for slow feeding and binge feeding from an example mouse. (B) Cumulative pump events for slow feeding and binge feeding from an example mouse (same data as in A). (C) Same as in (B) but for all aPC CaMK2⁺ mice (n= 481 for slow feeding trials and 241 trials for binge feeding trials from 8 mice). (D) Quantification of cumulative pump events within 15 s after feeding bout onset. Unpaired Student's t-test (n is the same as in C). (E) Lick events for slow feeding and binge feeding from an example mouse. (F) Cumulative lick events for slow feeding and binge feeding from an example mouse (same data as in E). (G) Same as in (F) but for all aPC CaMK2⁺ mice (n is the same as in C). (H) Quantification of cumulative lick events in 4 s after the onset of feeding bouts. Unpaired Student's t-test (n is the same as in C).

For (B) (C) (F) (G) data are shown as mean \pm s.e.m. For the box plot in (D) (H) the center line shows the median, the box limits show the quartiles, the whiskers show 1.5x the interquartile range, and the points show the outliers.

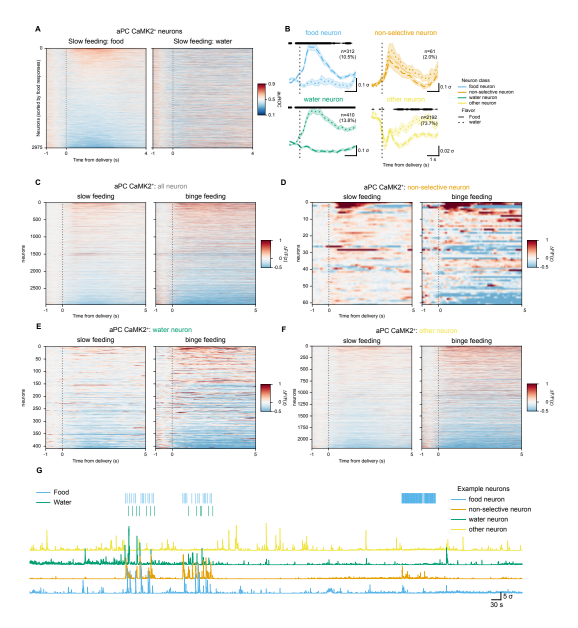


Figure S2. aPC CaMK2⁺ neuronal responses upon slow feeding and binge feeding.

(A) Area Under the Receiver Operating Characteristics (auROC) curve of aPC CaMK2⁺ neurons during slow feeding (see **Methods**, n is 2975 cells from 8 mice). (B) Slow feeding responses of aPC CaMK2⁺ neuronal subclasses for food and water deliveries (n=312 food neurons, 61 non-selective neurons, 410 water neurons, 2192 other neurons from 8 mice). The shaded line above denotes the adjusted P-values (Q-values) of each time point, with different line widths representing different values (from thin to thick: q<0.05, <0.01, <0.001). (C) Responses of individual aPC CaMK2⁺ neurons upon slow feeding and binge feeding (n is the same as in A). (D) Same as in (C) but with non-selective consumption neurons (n=61 cells from 8 mice). (E) Same as in (C) but with water-activated neurons (n=410 cells from 8 mice). (F) Same as in (C) but with non-responding neurons (n=2192 cells from 8 mice). (G) Example traces of aPC CaMK2⁺ neurons from a recording session.

For (**B**) data are shown as mean \pm s.e.m.

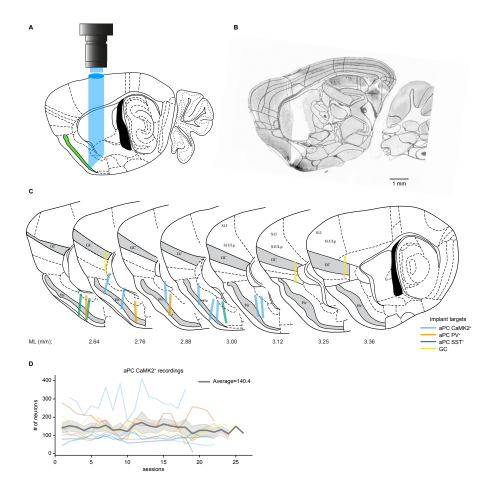


Figure S3. GRIN lens-Prism Implant coordinates and extracted cell numbers throughout experimental sessions.

(A) Schematic of implant coordination in the aPC. (B) GRIN lens-Prism path and GCaMP6f expression from an example mouse. (C) Reconstructed GRIN len-Prism coordinates in different cell types in the aPC (Fig.1, Fig.3) and GC (Fig.2). (D) Numbers of aPC CaMK2⁺ neurons extracted with *CNMFe* on each recording session (n= 8 mice).

For **(D)** data are shown as mean \pm s.e.m.

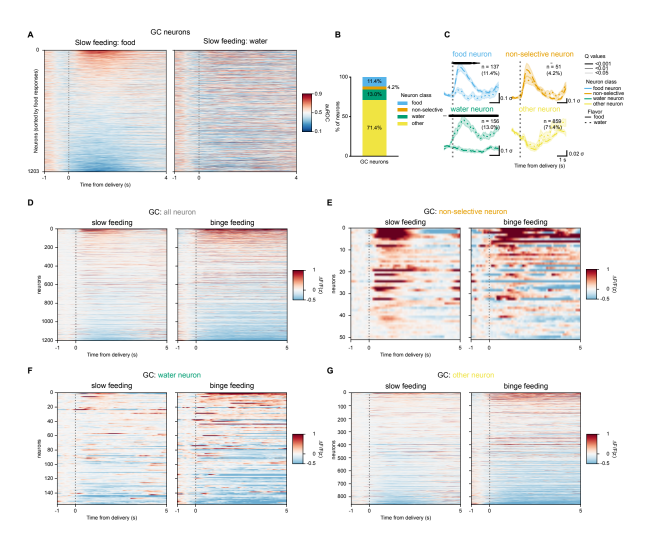


Figure S4. GC neuronal responses upon slow feeding and binge feeding.

(A) Area Under the Receiver Operating Characteristics (auROC) curve of GC neurons during slow feeding (see **Methods**). (B) Proportion of GC subclasses (n=1203 all neurons, 137 food neurons, 51 non-selective neurons, 156 water neurons, and 859 other neurons from 3 mice). (C) Food responses of subclasses of GC neurons during slow feeding for food and water deliveries (n is the same as in B). The shaded line above denotes the adjusted P-values (Q-values) of each time point, with different line widths representing different values (from thin to thick: q <0.05, <0.01, <0.001). (D) Responses of all GC neurons upon slow feeding and binge feeding (n=1203 cells from 3 mice). (E) Same as in (D) but with non-selective consumption neurons (n=51 cells from 3 mice). (F) Same as in (D) but with water-activated neurons (n=156 cells from 3 mice). (G) Same as in (D) but with non-responding neurons (n=859 cells from 3 mice).

For (C) data are shown as mean \pm s.e.m.

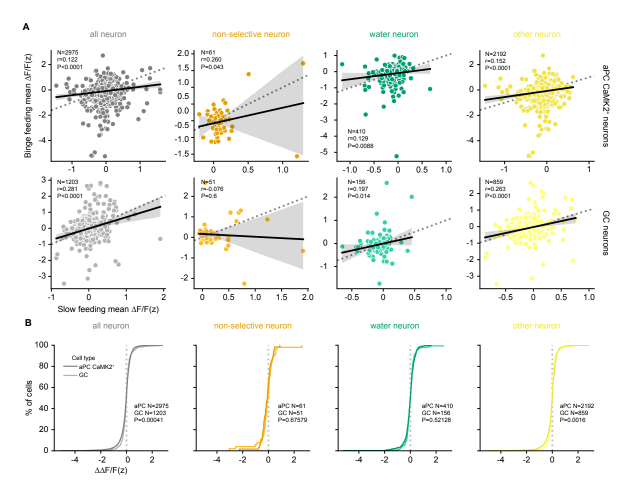


Figure S5. Distinct binge feeding-induced modulation in the aPC and the GC.

(A) Cell-wise comparison of neuronal responses upon slow feeding and binge feeding (For aPC CaMK2⁺ neurons, n=2975 for all neurons, 61 for non-selective neurons, 410 for water neurons, 2192 for other neurons from 8 mice. For GC neurons, n=1203 for all neurons, 51 for non-selective neurons, 156 for water neurons, and 859 for other neurons from 3 mice). (B) Cumulative distribution of the difference for each cell upon slow and binge feeding (binge feeding - slow feeding, $\Delta\Delta$ F/F(z), n is the same as in A. In (A) r and P represent the correlation coefficient and P-value of Pearson's r.

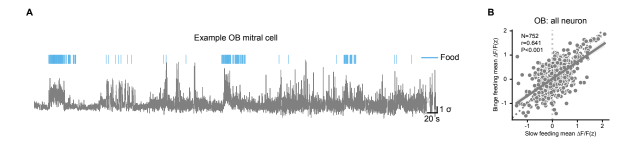


Figure S6. OB mitral cell example trace and cell-wise comparison.

(A) An example trace from an OB mitral cell upon slow feeding and binge feeding. (B) Cell-wise comparison of neuronal responses upon slow feeding and binge feeding (n=752 cells from 4 mice). In (B) r and P represent the correlation coefficient and P-value of Pearson's r.

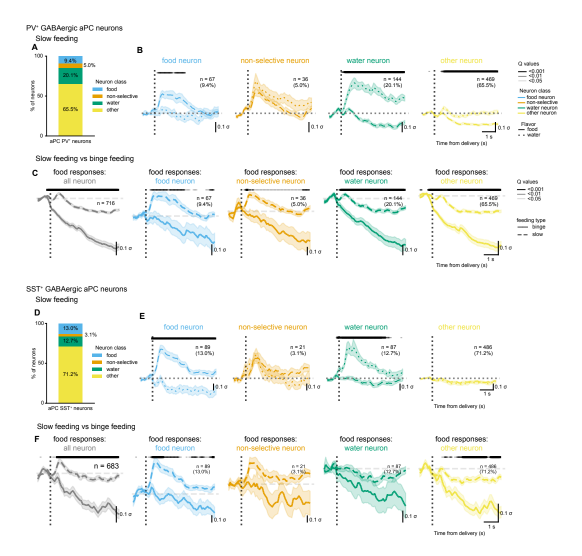


Figure S7. aPC GABAergic neuronal responses to slow feeding and binge feeding.

(A) Proportion of aPC PV $^+$ subclasses (n=67 cells for food neurons, 36 cells for non-selective neurons, 144 cells for water neurons, 469 cells for other neurons from 3 mice). (B) Responses of subclasses of aPC PV $^+$ neurons during slow feeding for food and water deliveries (n is the same as in a, except n= 716 cells for all neurons). (C) Food responses aPC PV $^+$ neuron subclasses during slow feeding and binge feeding (n is the same as in B). (D) Same as in (A) but for aPC SST $^+$ neurons (n=89 cells for food neurons, 21 cells for non-selective neurons, 87 cells for water neurons, 486 cells for other neurons from 3 mice). (E) Same as in (B) but for aPC SST $^+$ neurons (n is the same as in (C) but for aPC SST $^+$ neurons (n is the same as in (E) except n=683 cells in all neurons). The shaded line above denotes the adjusted P-values (Q-values) of each time point, with different line widths representing different values (from thin to thick: Q <0.05, <0.01, <0.001).

For **(B) (C) (E) (F)** data are shown as mean \pm s.e.m.

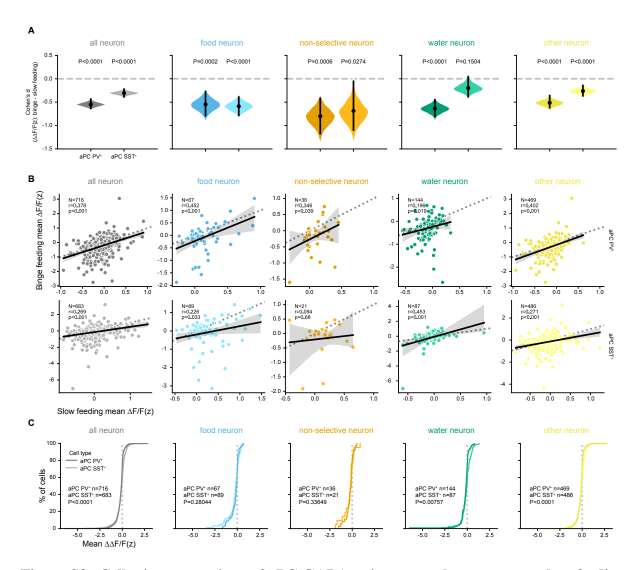


Figure S8. Cell-wise comparison of aPC GABAergic neuronal responses to slow feeding and binge feeding.

(A) Estimated effect size of binge feeding-induced suppression in aPC PV $^+$ and aPC SST $^+$ subclasses (For aPC PV $^+$ neurons, n=716 cells for all neurons, 67 cells for food neurons, 36 cells for non-selective neurons, 144 cells for water neurons, 469 cells for other neurons from 3 mice. For aPC SST $^+$ neurons, n=683 cells for all neurons, n=89 cells for food neurons, 21 cells for non-selective neurons, 87 cells for water neurons, 486 cells for other neurons from 3 mice). (B) Cell-wise comparison of neuronal responses upon slow feeding and binge feeding in aPC PV $^+$ and aPC SST $^+$ subclasses (n is the same as in A). (C) Cumulative distribution of the difference for each cell upon slow and binge feeding (binge feeding - slow feeding, $\Delta\Delta$ F/F(z)) in aPC PV $^+$ and aPC SST $^+$ subclasses (n is the same as in A).

For (B) r and P represent the correlation coefficient and P-value of Pearson's r.

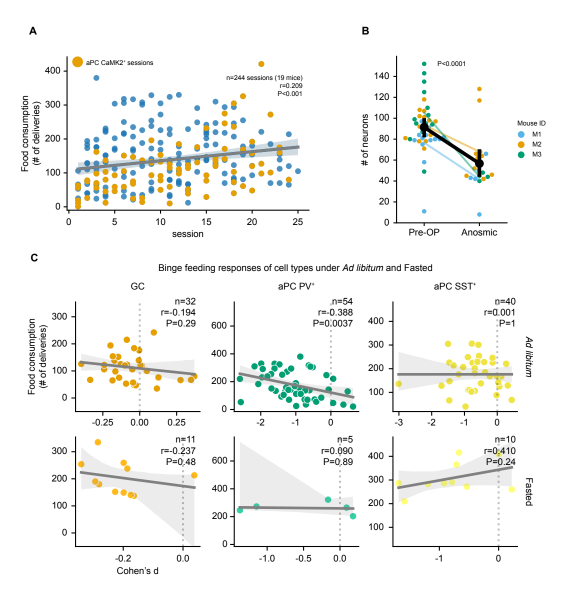


Figure S9. Effects of temporal progression, anosmia, and fasting.

(A) Food consumption (number of food deliveries) on subsequent recording sessions (n=244 sessions from 19 mice). Orange dots are mice with recordings in aPC CaMK2⁺ cells (n=84 sessions from 8 mice), blue dots represent all other mice. (B) Number of extracted cells in aPC CaMK2⁺ mice before and after Triton-X100 application (n=21 Pre-OP sessions and 15 anosmic sessions from the same 3 mice). Colors represent recordings from each mouse. c, Neuronal responses to binge feeding in the GC, aPC PV⁺, and aPC SST⁺under *ad libitum* (upper row) and fasting (lower row) conditions plotted against consumption (For GC, n=32 *ad libitum* sessions and 11 fasted sessions from 3 mice. For aPC PV⁺, n=54 *ad libitum* sessions and 10 fasted sessions from 3 mice.).

For **(B)** data are shown as mean \pm s.e.m. For **(A) (C)** r and P represent the correlation coefficient and P-value of Pearson's r.

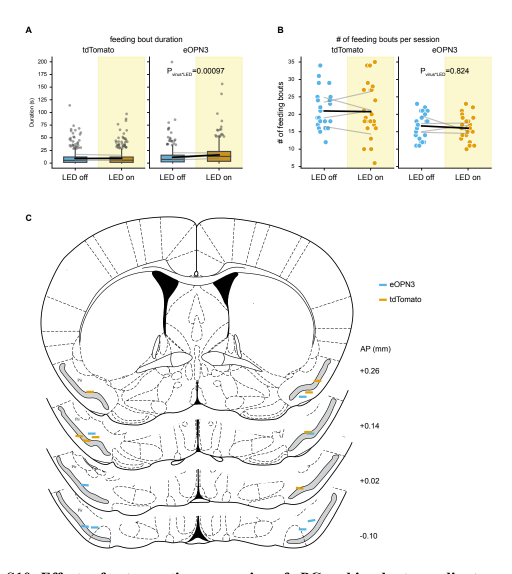


Figure S10. Effects of optogenetic suppression of aPC and implant coordinates.

(A) Effects of light stimulation on the duration of individual feeding bouts in tdTomato- and eOPN3-expressing mice (n=569 feeding bouts in LED off sessions and 562 feeding bouts in LED on sessions from 4 tdTomato mice. n=431 feeding bouts in LED off sessions and 425 feeding bouts in LED on sessions from 4 eOPN3 mice). (B) Effects of light stimulation on the number of feeding bouts per experimental session in tdTomato- and eOPN3-expressing mice (n=24 LED off sessions and 24 LED on sessions from 4 tdTomato mice and n=24 LED off sessions and 24 LED on sessions from 4 eOPN3 mice). Grey lines denote the data from the same mice, and the black line denotes the overall mean of the data. The p values are calculated from a linear mixed model. (C) Implant coordinates of optical fibers.

For the box plot in a and b, the center line shows the median, the box limits show the quartiles, the whiskers show 1.5x the interquartile range, and the points show the outliers.

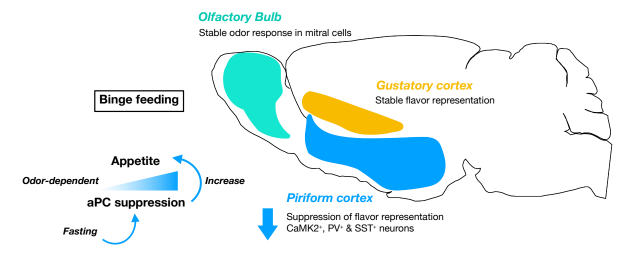
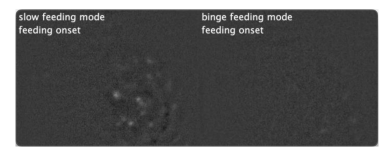


Figure S11. Graphic summary.

In this study, we found a feeding rate-dependent suppression of the olfactory flavor representation (Fig. 1 and Fig. 3) whereas the gustatory flavor representation is not affected by the feeding rate (Fig. 2). Olfactory inputs from the olfactory bulb remain stable across feeding rates (Fig. 3). We found the magnitude of binge feeding-induced aPC suppression correlates with appetite and the correlation depends on olfactory perception and metabolic state (Fig. 4). We further showed optogenetically suppression in the aPC upon feeding promotes appetite (Fig. 5).

SUPPLEMENTARY VIDEOS



Video. S1: Example Ca²⁺ video during slow feeding and binge feeding.

Left panel: Ca²⁺ transients of aPC CaMK2⁺ neurons during slow feeding. Right panel: Ca²⁺ transients of aPC CaMK2⁺ neurons during binge feeding from the same recording, the same mice.

SUPPLEMENTARY TABLES

Coordinations for injection and implantation

Table S1: Coordinates for viral injection and GRIN lens and prism implantation

Brain regions	Coordinates	Notes
	(AP/ML/DV, mm)	
aPC	0.32/-3.1/5.2	For viral injection
	0.32/-2.7/5.5	For implantation we used the bottom right corner of
		the prism of GRIN lens
GC	0.26/-3.6/4.0	For viral injection
	0.26/-3.1/4.1	For implantation we used the bottom right corner of
		the prism of GRIN lens
OB	4.3-4.6/±0.6 /0.3	For viral injection

Virus and construct

Table S2: Virus and construct table

Reagent or resource	Source	Identifier (RRID)
AAV1	Virus: Addgene	100833-AAV1
pAAV.Syn.Flex.GCaMP6f.WPRE.SV40		RRID:Addgene_100833
AAV8	Plasmid: Charité viral	Charité viral core id:
pAAV-Syn-iCre-RFP	core	BA-48c
	Virus: Charité viral	
	core	
AAV1	Virus: Addgene	104487-AAV1
pGP-AAV-syn-jGCaMP7s-WPRE		RRID:Addgene_104487
AAV9	Plasmid: Charité viral	Charité viral core id:
pAAV-hSyn-Flex-OPN3-mScarlet-	core	BA-575b
minWPRE	Virus: Charité viral	
	core	
AAV9	Plasmid: Charité viral	Charité viral core id:
hSyn-flox-tdTomato	core	BA-234a
	Virus: Charité viral	
	core	

734 Mouse line

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Table S3: Mouse line table

Reagent or resource	Source	Identifier (RRID)
C57BL/6N	Charité Central Animal	N/A
	Facility	
Ai93(TITL-GCaMP6f)-D	JAX	RRID:IMSR_JAX:024103
Rosa-tTA	JAX	RRID:IMSR_JAX:011008
Ai148(TIT2L-GC6f-ICL-tTA2)-D	JAX	RRID:IMSR_JAX:030328
CaMKII-CreT29	JAX	RRID:IMSR_JAX:005359
PV-Cre	JAX	RRID:IMSR_JAX:008069
SST-Cre	JAX	RRID:IMSR_JAX:013044

Software

Table S4: Software table

Reagent or re-	Source	Identifier (RRID/Version/reference)
source		
FIJI/ ImageJ		RRID:SCR_002285
		Version: 2.14.0/1.54f (Schindelin et al. 2012)
Python	Python	RRID:SCR_008394
	Software	Version: 3.9 (Rossum and Boer 1991)
	Founda-	
	tion	
Numpy		NumPy, RRID:SCR_008633
		Version: 1.21.5 (Harris et al. 2020)
Matplotlib		MatPlotLib, RRID:SCR_008624
		Version: 3.5.1 (Hunter 2007)
Seaborn		seaborn, RRID:SCR_018132
		0.12.2 (Waskom 2021)
Scikit-learn		scikit-learn, RRID:SCR_002577
		Version: 1.1.1 (Pedregosa et al. 2011)
Statsmodels		statsmodel, RRID:SCR 016074
		Version: 0.13.5 (Seabold and Perktold 2010)
Scipy		SciPy, RRID:SCR_008058
		Version: 1.8.0 (Virtanen et al. 2020)
Dabest		Version: 0.3.1 (Ho et al. 2019)
CNMFe/		Calcium Imaging data Analysis (Zhou et al. 2018; Giovan-
CaImAn		nucci et al. 2019), RRID:SCR_021533
		Git forked version:
		https://github.com/flatironinstitute/CaImAn.git@
		7dc5b42ab06c6a6b86ff1520dfc5b2334f335a78
Inscopix CN-		https://github.com/inscopix/isx-cnmfe-wrapper@v1.2
MFe wrapper		
Inscopix CN-	Inscopix	https://github.com/inscopix/inscopix-cnmfe
MFe		
Inscopix	Inscopix	
python API		
Inscopix Data	Inscopix	Version: 1.31, 1.6.0, 1.8.0
Processing		
Software		

Inscopix Data	Inscopix	Version: 1.31
Acquisition		
Software		
Bonsai		Bonsai, RRID:SCR_021512 (Lopes et al. 2015)
		Version: 2.4.0
FlyCapture2		Version: 2.11.3.425
Pyanpple		Version: 0.3.1 (Viejo et al. 2023)
LSMAQ		https://github.com/danionella/lsmaq
Suite2P		https://github.com/MouseLand/suite2p
		RRID:SCR_016434 (Pachitariu et al. 2016)
StichIt		https://github.com/SainsburyWellcomeCentre/StitchIt
BrainReg		https://github.com/brainglobe/brainreg (Claudi et al. 2020)
Cellfinder		https://github.com/brainglobe/cellfinder (Tyson et al. 2021)
BackingTray		https://github.com/SainsburyWellcomeCentre/BakingTray
R		Version: 4.2.2
		RRID:SCR_001905 (R Core Team 2021)
Rstudio		Version: 2022.12.0+353
		RRID:SCR_000432 (Posit team 2022)
lme4		Version: 1.1.31
		RRID:SCR_015654 (Bates et al. 2015)
Illustrator	Adobe	Version: 27.4.1, 2023
		RRID:SCR_010279

737 Statistical summary

Table S5: Statistical summary

				_	
Figure		Sample size			
Panel	description	cell/trial/session	mouse	statistical test	statical values
2e	aPC binge-slow all neuron Ensure neuron non-selective neuron water neuron other neuron	cell: 2975 312 61 410 2192	8	non-parametric two-sided approximate permutation t-test	P values: P<0.0001 P<0.0001 P=0.0132 P=0.0596 P<0.0001
2e	GC binge-slow all neuron Ensure neuron non-selective neuron water neuron other neuron	cell: 1203 137 51 156 859		non-parametric two-sided approximate permutation t-test	P values: P=0.4214 P=0.5394 P=0.34 P=0.368 P=0.2564
2f	aPC food neuron correlation	cell: 312	8	Pearson's correlation coefficent	r=-0.06479407831039309 P=0.25382944269503377
	GC food neuron	cell:		Pearson's correlation	r=0.43325819768924756
2f	correlation Cumulative distribution of Δzs- ΔF/F of aPC and GC	137		coefficent	P=1.2354901078156077e-07 KS_stats = 0.18793280928317424
2g	Ensure neurons cell types: aPC CaMK2 (same as 2e) aPC PV aPC SST GC (same as 2e)	312/137 cell: 2975 684 675 1203	8 3	Non-parametric two-sided approximate permutation t-	P=0.0020045609493748295 P values: P<0.0001 P<0.0001 P=0.0001 P=0.4214
<u>3m</u>	OB aPC CaMK2	752	4	test Linear Mixed Model: contribution of Cohen's d on Ensure consumption Linear Mixed Model: Fixed effects: intercept, recording session, Cohen's d Random effects: Mouse ID	P=0.1646 x2(1) = 2.0315 P=0.1541
4b	slow feeding vs Ensure consumption	sessions: 103	8	Testing for contribution of Cohen's d on Ensure intake Linear Mixed Model:	βinteraction = -36.306 ± 25.239 (standard errors)
				contribution of Cohen's d on Ensure consumption Linear Mixed Model: Fixed effects: intercept, recording session, Cohen's d Random effects: Mouse ID	χ2(1) = 11.546 P=0.0006791
4b	aPC CaMK2 binge feeding vs Ensure consumption	sessions:	۵	Testing for contribution of Cohen's d on Ensure intake	βinteraction = -116.462 ± 32.730 (standard errors)
4e	Buried food test Treatment: PBS Triton	10 3	10	Independent t-test	t=-5.140378686166757 P=0.000323044677564837
4f	aPC CaMK2 Pre-OP Cohen's d vs Ensure consumption	sessions:	3 (same mice)	Pearson's correlation coefficent	r=-0.504 P=0.02
4f	aPC CaMK2 Anosmic Cohen's d vs Ensure consumption aPC CaMK2 Cohen's d in Pre-Op	sessions: 15 sessions:	3 (same mice)	Pearson's correlation coefficent	r=-0.210 P=0.45 T = 0.49904460491871644
4g 4j	and Anosmic aPC CaMK2 Ad libitum Cohen's d vs Ensure consumption	21/15 sessions: 84		Pearson's correlation coefficent	P=0.6209617193931751 r=-0.43457505219920095 P=3.620129230550653e-05

Parent description Cell/mail/session mouse statistical test statical values	Fia		0		•	
aPC CaMK2	Figure Panel	description	Sample size cell/trial/session	mouse	statistical test	statical values
PC CaMM2		aPC CaMK2 Fasted Cohen's d vs	sessions:	5 (in 3 mice fasting protocol was not	Pearson's correlation	r=0.11785311537277354
Contents of in Ad Sessions 3/3 Independent I-lest P=0.007206291915825609		aPC CaMK2 Cohen's d in Ad	sessions:	, ,		T=2.7569856211682433
Private effects: Intercept, recording session, body weight, baseline feeding lime, sex, virus type, LED state Random effects: Mouse ID X2(LED state Random effects: X2(LED state Rando	4k	Cohen's d in Ad		3/3	Independent t-test	
APC optogenetic suppression vs					Fixed effects: intercept, recording session, body weight, baseline feeding time, sex, virus type, LED state, virus type*LED	
Ensure Intake						v2(1) = 11 631
Linear Mixed Model: Fixed effects: Intercept, recording session, body weight, baseline feeding time, sex, virus type, LED state, virus type LED state Random effects: Mouse ID X2(1) = 5.1898 P=0.02272 Testing for contribution of interaction of virus type and light stimulation on Feeding duration 4 durati	5a	Ensure intake tdTomato	24(LED off)/24(LED on) 24(LED off)/24(LED on)		interaction of virus type and light stimulation on Ensure	P=0.0006488 βinteraction = 70.7777 ± 20.0774
aPC optogenetic suppression vs Feeding duration tdTomato eOPN3	•				Fixed effects: intercept, recording session, body weight, baseline feeding time, sex, virus type, LED state, virus type*LED	
Cumulative pump/Ensure deliveries trials: Slow feeding	5h	suppression vs Feeding duration tdTomato	24(LED off)/24(LED on) 24(LED off)/24(LED on)		Mouse ID Testing for contribution of interaction of virus type and light stimulation on Feeding	P=0.02272 βinteraction = 47.23888 ± 20.43307
Cumulative pump/Ensure deliveries trials: Slow feeding						
pump/Ensure deliveries trials: Slow feeding 481 8 Independent t-test F=4.041159928317131 F=5.8911362087873396e-05	Supple	mentary figures				
events	S1d	pump/Ensure deliveries Slow feeding Binge feeding	481	8	Independent t-test	
S5a neuron correlation 2975 8 coefficent P=2.4028371776727368e-11	S1h	Slow feeding	481	8	Independent t-test	
aPC CaMK2 non-selective neuron correlation Fearson's correlation Fearson	S5a			8		
S5a neuron correlation 410 8 coefficent P=0.00879802235411608 B PC CaMK2 other neuron correlation cell: 21925 Pearson's correlation coefficent r=0.15224409159665298 GC all neuron correlation cell: 200.811566517498387 P=7.743112781578339e-13 S5a correlation r=0.2811566517498387 GC non-selective neuron correlation cell: 200.7575136213765701 Fearson's correlation neuron correlation 51 3 coefficent P=0.5972760763059162 GC water neuron correlation cell: 200.7575136213765701 Pearson's correlation r=0.1966433095696567 S5a correlation 156 3 coefficent p=0.013868121679462934 GC other neuron cell: 300.7575136213765701 P=0.013868121679462934 P=0.013868121679462934	S5a	selective neuron		8		r=0.2600348021314155
APC CaMK2 other neuron correlation 21925 8 Pearson's correlation r=0.15224409159665298 P=7.743112781578339e-13 GC all neuron correlation 1203 3 Pearson's correlation r=0.2811566517498387 P=2.71041468446213e-23 GC non-selective Cell: Pearson's correlation r=0.07575136213765701 S5a GC water neuron Cell: Pearson's correlation r=0.5972760763059162 GC water neuron Cell: Pearson's correlation r=0.15966433095696567 GC water neuron Cell: Pearson's correlation r=0.15966433095696567 GC other neuron Cell: Pearson's correlation r=0.263453681246707 GC other neuron Cell: Pearson's correlation r=0.2634536812426707 Cell: Pearson's correlation r=0.	S5a			8		
GC all neuron correlation 1203 3 Pearson's correlation r=0.2811566517498387 P=2.71041468446213e-23 GC non-selective Cell: Pearson's correlation r=0.07575136213765701 S5a neuron correlation 51 3 coefficent P=0.5972760763059162 GC water neuron Cell: Pearson's correlation r=0.19666433095696567 S5a Correlation 156 3 coefficent P=0.013868121679462934 GC other neuron Cell: Pearson's correlation r=0.2634536812426707 Cell: Pearson's co		aPC CaMK2 other	cell:		Pearson's correlation	r=0.15224409159665298
S5a neuron correlation 51 3 coefficent P=0.5972760763059162 GC water neuron correlation cell: pearson's correlation r=0.19666433095696567 r=0.19666433095696567 S5a GC other neuron 156 3 coefficent P=0.013868121679462934 GC other neuron cell: Pearson's correlation r=0.2634536812426707		GC all neuron	cell:	3	Pearson's correlation	r=0.2811566517498387
GC water neuron cell: Pearson's correlation r=0.19666433095696567 r=0.013868121679462934 GC other neuron cell: Pearson's correlation r=0.2634536812426707 r=0.2634536812676767 r=0.26345676767676767676767676			cell:	3	Pearson's correlation	r=-0.07575136213765701
GC other neuron cell: Pearson's correlation r=0.2634536812426707		GC water neuron	cell:		Pearson's correlation	r=0.19666433095696567
		GC other neuron	cell:		Pearson's correlation	r=0.2634536812426707

Figure		Sample size		-	
	description	cell/trial/session	mouse	statistical test	statical values
	ΔΔF/F distribution between aPC CaMK2 and GC	cells:	aPC=8	oranomean root	KS_stats=0.07012887948196735
S5b	all neuron	GC=1203		Kolmogorov-Smirnov test	P=0.00041157759310930927
S5b	ΔΔF/F distribution between aPC CaMK2 and GC non-selective neuron	cells: aPC=61 GC=51	aPC=8 GC=3	Kolmogorov-Smirnov test	KS_stats=0.1295403407264545 P=0.6757897458496079
	$\Delta\Delta$ F/F distribution between aPC CaMK2 and GC	cells: aPC=410	aPC=8		KS_stats=0.074859287054409
S5b	water neuron	GC=156	GC=3	Kolmogorov-Smirnov test	P=0.5212789564789919
S5b	ΔΔF/F distribution between aPC CaMK2 and GC other neuron	cells: aPC=2192 GC=859	aPC=8 GC=3	Kolmogorov-Smirnov test	KS_stats=0.07564442187911594 P=0.0016025784462102555
S6b	OB all neuron correlation	cells: 752	4	Pearson's correlation coefficent	r=0.6411402389745184 P=2.709628199058605e-88
S8a	aPC PV binge-slow all neuron Ensure neuron non-selective neuron water neuron other neuron	cell: 716 67 36 144 469		non-parametric two-sided approximate permutation t-test	F-2.709026199036003e-68 P values: P<0.0001 P=0.0002 P=0.0006 P<0.0001 P<0.0001 P<0.0001 P<0.0001 P<0.0001
S8a	aPC SST binge-slow all neuron Ensure neuron non-selective neuron water neuron other neuron	cell: 683 89 21 87 486	3	non-parametric two-sided approximate permutation t-test	P values: P<0.0001 P<0.0001 P=0.0274 P=0.1504 P<0.0001
S8b	aPC PV all neuron correlation	cell: 716	3	Pearson's correlation coefficent	r=0.3784480776726223 P=8.456499073277892e-26
S8b	aPC PV food neuron correlation	cell: 67	3	Pearson's correlation coefficent	r=0.45177311835390116 P=0.0001242739664864955
S8b	aPC PV non-selective neuron correlation	cell:		Pearson's correlation coefficent	r=0.3460256619145978 P=0.038712553293138004
S8b	aPC PV water neuron correlation	cell: 144	3	Pearson's correlation coefficent	r=0.19592988760587732 P=0.018596309032818777
S8b	aPC PV other neuron correlation	cell: 469	3	Pearson's correlation coefficent	r=0.40234957084687484 P=1.1198215205307785e-19
S8b	aPC SST all neuron correlation	cell: 683	3	Pearson's correlation coefficent	r=0.2693313778561128 P=8.151681475471994e-13
S8b	aPC SST food neuron correlation	cell: 89	3	Pearson's correlation coefficent	r=0.22598666216076777 P=0.03321740625273849
S8b	aPC SST non- selective neuron correlation	cell: 21	3	Pearson's correlation coefficent	r=0.09443509682038038 P=0.6838838663964277
S8b	aPC SST water neuron correlation	cell: 87	3	Pearson's correlation coefficent	r=0.4534392424767807 P=1.0313185179901372e-05
S8b	aPC SST other neuron correlation	cell: 486		Pearson's correlation coefficent	r=0.2705659219270069 P=1.3375421968087283e-09
S8c	∆∆F/F distribution between aPC PV and aPC SST all neuron	cell: aPC_PV=716 aPC_SST=683	aPC_PV=3	Kolmogorov-Smirnov test	KS_stats=0.17635799995092305 P=5.550674405045618e-10
S8c	$\Delta\Delta$ F/F distribution between aPC PV and aPC SST food neuron	cell: aPC_PV=67 aPC_SST=89	aPC_PV=3	Kolmogorov-Smirnov test	KS_stats=0.15495555928224047 P=0.2804364616666969
S8c	ΔΔF/F distribution between aPC PV and aPC SST non-selective neuron	cell: aPC_PV=36 aPC_SST=21	aPC_PV=3 aPC_SST=3	Kolmogorov-Smirnov test	KS_stats=0.24603174603174602 P=0.3364929433449463
S8c	∆∆F/F distribution between aPC PV and aPC SST water neuron	cell: aPC_PV=144 aPC_SST=87	aPC_PV=3 aPC_SST=3	Kolmogorov-Smirnov test	KS_stats=0.22246168582375478 P=0.007573604429318253
S8c	$\Delta\Delta$ F/F distribution between aPC PV and aPC SST other neuron	cell: aPC_PV=469 aPC_SST=486	aPC_PV=3 aPC_SST=3	Kolmogorov-Smirnov test	KS_stats=0.19500820412926548 P=1.973926024904841e-08

Figure		Sample size		-	
Panel	description	cell/trial/session	mouse	statistical test	statical values
	Food consumption in different recording	sessions:		Pearson's correlation	r=0.2094528185645767
S9a	sessions Neuron numbers after anosmia	all sessions=244 sessions:	all mice = 19	coefficent	P=0.000996393855726413
S9b	Pre-OP Anosmia	21 15	3 3	Independent t-test	t = 5.068140807075272 P=4.387642827378711e-06
S9c	Binge feeding responses of cell types in ad libitum in GC	sessions: 32	3	Pearson's correlation coefficent	r=-0.1938216819439083 P=0.2878184757849871
S9c	Binge feeding responses of cell types in ad libitum in aPC PV	sessions: 54	3	Pearson's correlation coefficent	r=-0.3884010496175349 P=0.003704862297625621
S9c	Binge feeding responses of cell types in ad libitum in aPC SST	sessions:	3	Pearson's correlation coefficent	r=0.0005763290584942845 P=0.9971839222546877
S9c	Binge feeding responses of cell types in fasted in GC	sessions:	3	Pearson's correlation coefficent	r=-0.23699667524513704 P=0.48288660842195086
S9c	Binge feeding responses of cell types in fasted in aPC PV	sessions: 5	3	Pearson's correlation coefficent	r=-0.08959548146743346 P=0.8860762962853914
S9c	Binge feeding responses of cell types in fasted in aPC SST	sessions: 10	3	Pearson's correlation coefficent	r=0.4095755900514694 P=0.23982700271970062
				Linear Mixed Model: Fixed effects: intercept, recording session, body weight, baseline feeding time, sex, virus type, LED state, virus type*LED state	
				Random effects: Mouse ID	
	aPC optogenetic suppression vs feeding bout duration	sessions: 24(LED off)/24(LED on)		Testing for contribution of the interaction of virus type and light stimulation on	x2(1) = 10.882 P=0.0009711
S10a	tdTomato eOPN3	24(LED off)/24(LED on) 6/6 sessions for each mouse	4	duration of individual feeding bout	βinteraction = 4.112 ± 1.245 (standard errors)
				Linear Mixed Model: Fixed effects: intercept, recording session, body weight, baseline feeding time, sex, virus type, LED state, virus type*LED state	
	aPC optogenetic suppression vs number of feeding	sessions:		Random effects: Mouse ID Testing for contribution of	χ2(1) = 0.0495 P=0.824
S10b	bout tdTomato eOPN3	24(LED off)/24(LED on) 24(LED off)/24(LED on) 6/6 sessions for each mouse		interaction of virus type and light stimulation on numbers of feeding bouts	βinteraction = -0.4337 ± 1.9494 (standard errors)