European Journal of Neuroscience, Vol. 35, pp. 146-159, 2012

doi:10.1111/j.1460-9568.2011.07916.x

BEHAVIORAL NEUROSCIENCE

Dopaminergic enhancement of local food-seeking is under global homeostatic control

Jeff A. Beeler, 1 Cristianne R. M. Frazier2 and Xiaoxi Zhuang1,2

¹Department of Neurobiology, The University of Chicago, 924 E 57th St. R222, Chicago, IL 60637, USA

Keywords: dopamine, dopamine transporter knock-down, feeding, homeostatic regulation, mouse, semi-naturalistic homecage operant

Abstract

Recent work has implicated dopaminergic mechanisms in overeating and obesity with some researchers suggesting parallels between the dopamine dysregulation associated with addiction and an analogous dysregulation in obesity. The precise role of dopamine in mediating reward and reinforcement, however, remains controversial. In contrast to drugs of abuse, pursuit of a natural reward, such as food, is regulated by homeostatic processes that putatively maintain a stable energy balance keeping unrestrained consumption and reward pursuit in check. Understanding how the reward system is constrained by or escapes homeostatic regulation is a critical question. The widespread use of food restriction to motivate animal subjects in behavior paradigms precludes investigation of this relationship as the homeostatic system is locked into deficit mode. In the present study, we examined the role of dopamine in modulating adaptive feeding behavior in semi-naturalistic homecage paradigms where mice earn all of their food from lever pressing. We compared consumption and meal patterning between hyperdopaminergic dopamine transporter knock-down and wild-type mice in two paradigms that introduce escalating costs for procuring food. We found that hyperdopaminergic mice exhibited similar demand elasticity, weight loss and energy balance in response to cost. However, the dopamine transporter knock-down mice showed clear differences in meal patterning. Consistent with expectations of enhanced motivation, elevated dopamine increased the meal size and reduced intrameal cost sensitivity. Nonetheless, this did not alter the overall energy balance. We conclude that elevated dopamine enhances the incentive or willingness to work locally within meals without shifting the energy balance, enhancing global food-seeking or generating an energy surplus.

Introduction

The cause of the dramatic rise in obesity in recent years (Wang & Beydoun, 2007; Center for Disease Control and Prevention, 2010) is not fully understood and no clear program for reversing this trend has yet emerged. Historically viewed primarily from the perspective of biologically determined homeostatic mechanisms, energy balance is increasingly seen as arising from complex interactions between genetic and environmental factors (Hill, 2006; Neel, 1999). The last decade has seen a growing appreciation of the role of non-homeostatic systems and processes, particularly the reward and incentive system and reinforcement learning (Kelley *et al.*, 2005a; Berthoud, 2007; Rowland *et al.*, 2008; Berridge *et al.*, 2010; Kenny, 2010).

Recent work has implicated dopaminergic mechanisms in overeating and obesity, although the nature of their role is unclear. Both increased and decreased reward sensitivity, in overweight and obese subjects, respectively, have been observed (Kenny, 2010), making simple attributions about the role of dopamine difficult. An idea with growing currency is to liken overeating and obesity to addiction, invoking dopaminergic dysregulation in the reward system as a key

mechanism underlying excessive caloric intake, lack of executive control and the resulting net positive energy balance and obesity (Volkow & Wise, 2005; Volkow et al., 2010). Notably, however, the role of dopamine in both addiction and natural reward remains controversial (Dayan & Balleine, 2002; Redish, 2004; Wise, 2004; Balleine, 2005; Everitt & Robbins, 2005; Berridge, 2007; Di Chiara & Bassareo, 2007; Salamone, 2007; Schultz, 2007). Viewing food as an addiction (broadly and widely construed as the 'hijacking' of the natural reward system) adds an additional layer of complexity to the already controversial role of dopamine in addiction and reward. Specifically, it suggests that natural rewards can 'hijack' the natural reward system, begging the question - what are the presumably homeostatic controls on this natural reward system and what causes them to fail? It is precisely the absence of such regulatory systems that is believed to underlie the vulnerability of the reward system to drugs of abuse (Di Chiara, 2005). Although fairly recent work has begun to highlight the interactions between reward/incentive and homeostatic mechanisms, much of this work is in its early stages and continues to evolve (Kelley et al., 2005b; Fulton et al., 2006; Hommel et al., 2006; Figlewicz et al., 2007; Palmiter, 2007; Berthoud & Morrison, 2008; Lutter & Nestler, 2009; Davis et al., 2010; Figlewicz & Sipols, 2010).

Behavioral studies of incentive and reward processes traditionally use food (or water) restriction to motivate subjects to perform the

Correspondence: Dr J. A. Beeler, as above. E-mail: jabeeler@uchicago.edu

Received 1 August 2011, revised 12 September 2011, accepted 19 September 2011

²Committee on Neurobiology, The University of Chicago, Chicago, IL, USA

experimental task. Doing so artificially locks homeostatic systems into a deficit state, effectively driving motivation but precluding investigation of how homeostatic and incentive processes jointly contribute to self-regulated energy balance behaviors. Such investigation remains an important challenge. Consequently, we have adopted a seminaturalistic homecage operant paradigm in which mice earn all of their food through lever pressing with no explicit food restriction, allowing both homeostatic and incentive processes to determine ongoing behavior.

In the present study, we examined the role of dopamine in adaptively modulating appetitive behavior in response to environmental cost contingencies. Using dopamine transporter knock-down (DATkd) mice that exhibit elevated extracellular dopamine and increased tonic firing of dopamine neurons (Zhuang et al., 2001), we investigated how hyperdopaminergia alters subjects' consummatory behavior in response to increasing costs. Associated with enhanced incentive (Berridge, 2007) and decreased sensitivity to cost (Salamone & Correa, 2002), we hypothesized that increased tonic dopamine would enhance adaptation to high costs associated with acquiring food. Increased dopamine did not substantially increase resistance to cost or improve survival. However, it did alter meal patterning in favor of longer but fewer meals. This finding is consistent with a role for dopamine in increasing incentive and/or decreasing cost sensitivity locally within individual meals, whereas the overall energy balance remains under homeostatic regulation. We propose that the dopaminemediated enhancement of incentive does not act globally to increase episodes of goal pursuit or food-seeking but is restricted locally to augmenting the pursuit of a goal once initiated.

Materials and methods

Subjects

Mice for the demand study were all male between 6 and 8 weeks of age. Mice in the homecage studies were all male (progressive ratio) or mixed sex (progressive interval) between 12 and 16 weeks of age. All mice were housed under standard 12:12 light cycles with free access to water and access to food as described below. Wild-type C57BL/6 mice were obtained from Jackson Laboratories. All animal procedures were approved by the Institutional Animal Care and Use Committee at The University of Chicago.

Dopamine transporter knock-down mice

The DATkd mice were from an established colony backcrossed with C57BL/6 mice for more than 10 generations. The DATkd mice have been described and extensively characterized (Zhuang et al., 2001; Pecina et al., 2003; Cagniard et al., 2006a,b; Yin et al., 2006; Beeler et al., 2010). These mice exhibit an 85% reduction in dopamine transporter (DAT) expression, resulting in elevated extracellular dopamine and increased tonic dopamine cell activity (Zhuang et al., 2001; Cagniard et al., 2006b). Phasic dopamine cell activity is unaltered with this mutation (Cagniard et al., 2006b); however, there is a 25% reduction in the amplitude of dopamine release arising from phasic activation (Zhuang et al., 2001). Unlike the DAT knock-out mice (Bosse et al., 1997), these mice show no developmental abnormalities and multiple studies have demonstrated no deficits in learning (Cagniard et al., 2006b; Yin et al., 2006; Beeler et al., 2010). High-performance liquid chromatography analysis of tissue dopamine comparing the DATkd with wild-type mice showed that intracellular dopamine was diminished and dopamine turnover, as reflected by 3,4dihydroxyphenylacetic acid (DOPAC): dopamine and homovanillic acid (HVA): dopamine ratios, was increased in the dorsal and ventral striatum as well as the hypothalamus, the greatest effect being observed in the dorsal striatum (Table 1). Within the prefrontal cortex, dopamine reuptake is mediated primarily by norepinephrine transporters and available evidence suggests that diminished DAT does not significantly alter the pharmacokinetics of dopamine reuptake in the prefrontal cortex (PFC) (Sesack et al., 1998; Mundorf et al., 2001; Moron et al., 2002).

Behavior setup and housing

Mice were singly housed in standard cages equipped with two levers placed on one side of the cage approximately 15 cm apart with a food hopper between the levers (Med-Associates, St Albans, VT, USA). A pellet dispenser delivered 20 mg grain-based precision pellets (Bio-Serv, Frenchtown, NJ, USA) contingent on lever presses according to a programmed schedule. No other food was available. Water was available ad libitum. Upon initial placement in the operant homecages, three pellets were placed in the food hopper and the first 50 lever presses on the active lever yielded a pellet (continuous reinforcement), after which the experimental design was initiated. All mice acquired the lever-pressing response overnight. One lever was active and yielded reward, the other was inactive and had no programmed consequences.

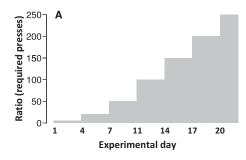
Behavioral paradigms

In the demand study, the active lever operated on a fixed-ratio (FR) schedule incremented across the experiment (5, 20, 50, 100, 150, 200, 250, 300). Each ratio was in effect for 3 days (Fig. 1A). Data were collected using MED-PC IV software (Med-Associates) and flushed daily at 09:00 h. The program tracked total daily consumption, active and inactive lever presses, the number of meals in a 24 h period, the average size and duration of meals, the average intermeal interval, and the average rate of pressing across meals (i.e. number of presses/meal duration). A meal was qualitatively defined as a sustained period of eating and effort. We operationally defined a meal as follows – the first pellet earned signaled the 'start' of a meal. A meal was considered as

TABLE 1. High-performance liquid chromatography analysis of intracellular dopamine and dopamine turnover (ng/mg) comparing wild-type (WT) and DATkd mice

	Dopamine		DOPAC : DA		HVA: DA	
	WT	DATkd	WT	DATkd	WT	DATkd
Dorsal striatum Ventral striatum Hypothalamus	86.83 ± 6.8 37.68 ± 1.1 1.84 ± 0.087	40.16 ± 7.4 26.45 ± 3.4 1.25 ± 0.134	0.064 ± 0.003 0.131 ± 0.008 0.259 ± 0.034	0.135 ± 0.017 0.146 ± 0.009 0.305 ± 0.021	0.109 ± 0.008 0.128 ± 0.014 0.572 ± 0.072	0.491 ± 0.006 0.278 ± 0.040 0.905 ± 0.098

All genotype differences significant, P < 0.01. DA, dopamine; DOPAC, 3,4-dihydroxyphenylacetic acid; HVA, homovanillic acid.



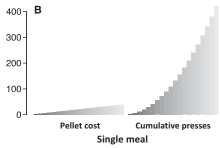


Fig. 1. Schematic of cost schedule for (A) homecage demand paradigm and (B) homecage progressive ratio paradigm.

terminated after 30 min without earning a reward, at which time the meal termination was recorded as having occurred 30 min previously (Chaney & Rowland, 2008). Single-pellet 'meals' were discarded as mice tend to almost continuously sample the levers at a low rate unless asleep. Such random pressing occasionally results in an earned pellet that does not reflect sustained effort or eating. An event recorder coded and time-stamped every lever press, pellet delivery, meal start and meal end. These data were used to construct a meal dataset (i.e. rather than daily averages across meals) for more detailed analysis.

In the homecage progressive ratio, mice were given 24 h of training in which the first 50 pellets were earned on an FR1 schedule, followed by FR5 until training was over. Following this, the progressive ratio (PR2) was initiated. After each reward during a meal, the lever press requirement for the next pellet was increased by 2. Although this resulted in only a moderate increase in the ratio requirement for each pellet, the cumulative number of presses required to continue a meal grew dramatically (Fig. 1B). The ratio reset back to 2 after 30 min of inactivity. A 1 s cue light signaled each pellet delivery. When the ratio reset, the cue light was set to constant illumination until the next active lever press initiating a new meal. The breakpoint was defined as the last successfully completed ratio. The program tracked the number, size and duration of meals (defined as a progressive ratio reset) as well as the average breakpoint, total consumption and total lever presses, both active and inactive. The progressive interval was identical (i.e. training, cue light) to the progressive ratio except for a delay interval rather than required ratio increments. In the progressive interval paradigm only one press was required to dispense a pellet; however, after each pellet delivery, there was a delay before another pellet became available. This delay progressively incremented analogous to the progressive ratio above and also reset after 30 min of inactivity.

Analyses

All statistical analyses were performed using R statistical software [R version 2.12.1 (2010-12-16); The R Foundation for Statistical Computing, http://www.r-project.org]. To analyze the mixed within-and between-groups design of the demand study (Figs 3–5), unbalanced due to mice dropping out at different points, we used a linear mixed-effects model (NLME package) with mouse as a random, within-subjects variable as follows:

$$Y = \text{initial body weight (where applicable)} + genotype \times cost, random = 1 | mouse.$$

Two-sample comparisons used *t*-tests and survival analysis was performed using the R survival package. Where data were balanced (e.g. Supporting Information Fig. S2), traditional repeated-measures ANOVA was used. The meal data (Figs 7 and 8; Table 1) were

constructed from the same dataset except that the unit of observation was individual meals rather than daily averages. Within the meal data, we removed meals consisting of a single pellet, considering these as arising from accumulating, random pressing rather than occurring within a bout of focused effort. There were no differences between genotypes in terms of the number of single-pellet meals overall (t = 0.5808, P = 0.571) during either the active (t = 0.6935,P = 0.499) or inactive (t = 0.3959, P = 0.698) cycle. This dataset was also unbalanced as mice varied in their daily number of meals. In addition, meal size and duration were periodic with a bimodal distribution corresponding to the active and inactive circadian phases. To model these, we used linear regression incorporating both fixed effects and covariance structures, specifically matrices for covariance by mouse (M), start time of meal (T) and mouse \times time (M*T). These analyses were performed using the R package Regress. To assess the contribution of individual factors, we removed each individually, starting with interaction terms, and compared each subsequent loglikelihood of the reduced model with the log-likelihood of the full model using chi-square distribution. When a term did not contribute to a significantly better model, it was removed (the least significant being removed first) and was then considered as the full model. Only two terms were removed, both from the intermeal duration model. Once the models with the fewest parameters that best fit the data were determined, P-values within that model were determined using zscores [i.e. pnorm(coefficient/SE of coefficient)].

To evaluate elasticity, we used an established model developed by Hursh & Silberberg (2008) as follows:

$$\log Q = \log Q_0 + k(e^{-\alpha P} - 1)$$

where Q is consumption/demand, Q_0 is baseline demand (we used FR5 as Q_0), k determines the range of the demand, P is price (ratio) and α represents the elasticity coefficient.

In Fig. 9, energy balance dynamics were evaluated as follows.

Peak energy stores occurred at the termination of a meal and trough energy stores at initiation. To approximate the energy balance thresholds for initiating and terminating meals, we defined the initiation threshold as the degree to which a previous energy peak is depleted prior to initiating a new meal:

prior meal size/ $(k_{\text{depletion}} \times \text{intermeal duration}) = \text{initiation threshold}$

where $k_{\rm depletion}$ is a constant that represents the average rate of basal energy expenditure, which we set equal to 1. This value represents the percentage of energy stores remaining since the previous peak such that larger numbers reflect less depletion, whereas smaller numbers reflect greater depletion. To make this more intuitive, we plotted (1– initiation threshold), which reflects the percentage to which energy previously ingested is depleted.

We assessed the termination thresholds as follows:

current meal size/ $(k_{\text{depletion}} \times (\text{intermeal duration} + \text{meal duration}))$ = termination threshold

again setting the constant $k_{\text{depletion}}$ equal to 1. This reflects the degree to which the current meal restores energy lost since the last energy peak. Larger numbers reflect greater energy restoration. These data were analyzed using a linear mixed-effects model (NLME package).

Results

Dopamine transporter knock-down mice exhibit no differences in consumption or body weight at baseline low-cost conditions

There were no initial differences between genotype in body weight (Fig. 2A; t = 1.55, P = 0.124, N = 15). Consistent with previous reports (Beeler et al., 2010), during the low-cost, FR5 portion of the demand experiment, there was no significant difference between genotypes in either consumption (Fig. 2B; genotype main, $F_{1,15} = 0.6857$, P = 0.4215) or body weight (genotype main,

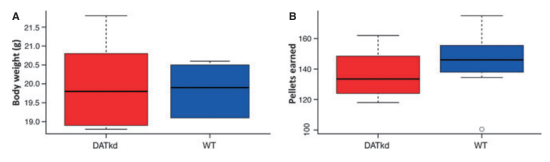


FIG. 2. Baseline comparison of genotypes. (A) Initial body weight and (B) daily consumption at low-cost FR5 schedule. N = 8. WT, wild-type.

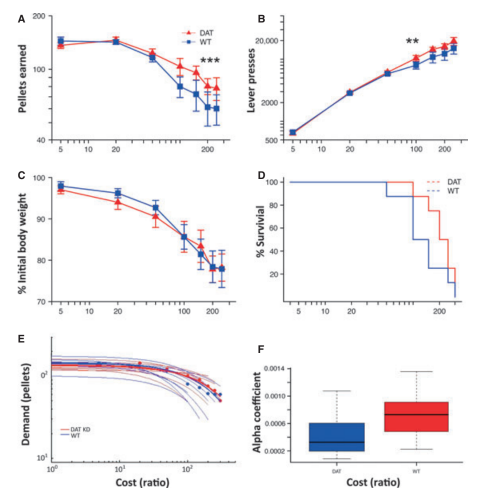


Fig. 3. Demand, survival and elasticity. (A) Average daily consumption of 20 mg pellets at each ratio/cost (cost × genotype, ***P < 0.001). (B) Average daily lever presses (cost × genotype, **P < 0.01). (C) Average daily body weight as a percentage of initial weight. (D) Survival curve (defined as asymptotic lever pressing). (E) Demand curve (see Materials and methods) of individual mice (light traces) with average for each genotype represented as points with bold trace. (F) Boxplot of elasticity coefficient for each genotype. N = 8. KD, knock-down; WT, wild-type.

 $F_{1,15} = 0.1377$, P = 0.7161, data not shown). These data indicate that the DATkd and wild-type mice exhibited similar energy balance under baseline conditions prior to environmental challenge.

Increased dopamine does not improve adaptation to escalating costs or survival

In the first experiment, mice were housed in homecages equipped with operant levers through which they were required to earn their entire food intake through lever pressing. The 'cost' of a 20 mg grain pellet changed on an incrementing schedule, starting at low cost (FR5) and increasing to high cost (FR250), allowing 3 days on each cost schedule. The change in consumption as cost increased was linear (see Supporting Information Fig. S1) and we used a linear mixed-effects model to evaluate statistical significance (see Materials and methods). As cost increased, both genotypes significantly decreased their daily consumption (Fig. 3A; cost main effect, t = 10.03, P < 0.0001), although increasing overall responding (Fig. 3B; cost main effect, t = 15.7, P < 0.0001). A significant genotype by cost interaction could be observed, indicating that at higher costs the DATkd mice

pressed and consumed more than the wild-type mice (Fig. 3A and B; genotype \times cost, pellets, t = 3.49, P = 0.0008; genotype \times cost, lever presses, t = 3.05, P = 0.0031), consistent with expectations that hyperdopaminergic mice would be more willing to work (Salamone et al., 1994, 1997). However, the magnitude of this effect, compared with the cost-related decline in responding, was surprisingly small. Presses on the inactive lever as a percentage of total presses were not significantly different by genotype (Supporting Information Fig. S2A; genotype main effect, t = 0.689, P = 0.5026, genotype × cost, t = 0.239, P = 0.8112), although these declined dramatically for both groups as cost increased (cost main effect, t = 5.04, P < 0.0001). The rate of lever pressing between genotypes was not significantly different (Supporting Information Fig. S2B; genotype main effect, $t = 0.380, P = 0.7098, \text{ genotype} \times \text{cost}, t = 1.13, P = 0.2597)$ and neither were there any significant differences in inter-response times or post-reinforcement pauses (Supporting Information Fig. S2C and D; inter-response times, genotype main t = 0.001, P = 0.9991; postreinforcement pauses, genotype main, t = 0.9234, P = 0.3714).

The minimal impact of this genotype difference was observed in the adaptive outcomes. Despite slightly more consumption at higher costs,

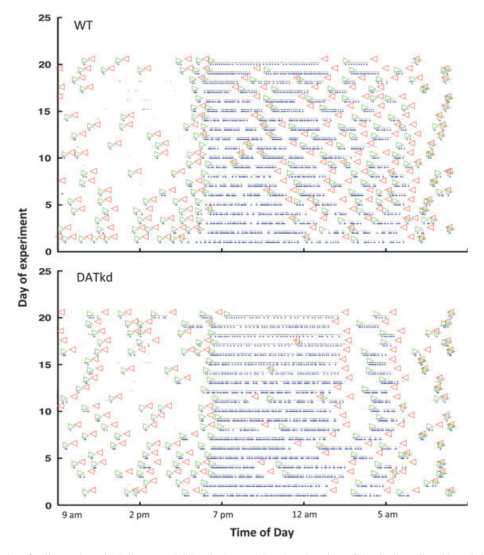


FIG. 4. Example raster plot of pellets and meals. Pellets earned (blue dots) were plotted against time of day horizontally with each line representing a single experimental day (vertical axis). Meal initiation as identified by the program (30 min without a pellet delivery) is marked with open green arrows and termination with open red arrows. One wild-type (WT) and one DATkd mouse are shown.

there was no resulting significant difference in body weight between the groups (Fig. 3C; genotype main effect, t = 0.377, P = 0.7122) or in survival, defined here as the breakpoint at which individual responding asymptotes regardless of increasing cost, generally necessitating removal from the experiment (Fig. 3D; survival difference chi-square, P = 0.179).

In (neuro)economics, the degree to which consumption or 'demand' adjusts in response to associated costs is referred to as elasticity. The present data suggest that although the hyperdopaminergic mice exhibited a slightly greater resistance to cost (i.e. slightly greater high-cost responding, Fig. 3A and B), overall they remained subject to the same processes that induce food- and energy-related elasticity in response to escalating costs. To look at the impact of hyperdopaminergia specifically on elasticity, we fitted the data to a wellestablished model developed by Hursh & Silberberg (2008) (see Materials and methods). We observed no difference between genotype in this model fit (Fig. 3E) and no statistically significant difference in alpha values, the parameter that measures elasticity (Fig. 3F; t = 1.509, P = 0.1599).

These results suggest that although dopamine does alter an animal's response to cost (Fig. 3A and B), it is not the only, or most important, factor mediating cost sensitivity. In particular, increased cost in this paradigm induces environmental scarcity, i.e. food is harder to come by as the experiment progresses. Consequently, scarcity may induce energy conservation, possibly through homeostatic mechanisms, which are either not significantly altered by dopaminergic function or actually modulate dopamine (Hommel et al., 2006; Lutter & Nestler, 2009; Figlewicz & Sipols, 2010). This would be consistent with the observed similarity in demand and survival between the genotypes.

It is possible that, at higher costs, the energy required to press the lever enough to earn a pellet is greater than the energy obtained, artificially forcing an energy deficit. This problem, however, would only apply to those mice that sustained pressing at higher costs and would not explain those that decreased demand at lower costs. It is difficult to assess the caloric costs of lever pressing; however, the results presented in the remainder of this article argue that the observed decrease in demand reflects a regulatory process rather than an artifact arising from forced energetic deficits. This issue will be revisited in the Discussion.

Increased dopamine alters meal patterning

To further investigate the impact of hyperdopaminergia on behavior in this paradigm, we analyzed meal patterning. Meals were defined as starting with the first pellet earned and terminating after 30 min having not received a pellet (subtracting 30 min from the meal duration). Figure 4 shows two example raster plots of meals across the course of the experiment. We analyzed the number of meals (Fig. 5A), meal size (Fig. 5B), meal duration (Fig. 5C) and length of intermeal intervals (Fig. 5D). Again, at low-cost baseline (FR5), no differences were observed between genotypes (Fig. 5A-D). However, with escalating costs, meal patterning diverged. Overall, the DATkd mice ate fewer (Fig. 5A; genotype main effect, t = 3.14, P = 0.0078) but larger meals with meal size, in contrast to wild-type mice, showing relative insensitivity to cost (Fig. 5B; genotype \times cost, t = 3.19, P = 0.0020). Larger meals were compensated by marginally significantly greater intermeal intervals (Fig. 5D; genotype main effect, t = 1.85, P =0.0861). These data suggest that increased dopamine, although having only a limited impact on overall energy balance in response to increasing cost and scarcity, nonetheless had a clear and highly significant effect on meal patterns. To determine more precisely what aspect of meal patterning was altered by increased dopamine (i.e. what component of behavior), we developed a model of the meal data.

Meal pattern model - incorporating time

We started with the basic premise that meal patterning is the composite of two processes - initiation and termination of meals. We

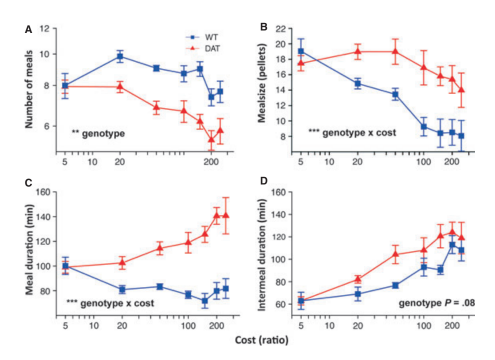


Fig. 5. Meal patterning. (A) Average number of meals per day across ratio/costs, (B) average meal size, (C) average duration of meals and (D) average intermeal duration. **P < 0.01, ***P < 0.001. N = 8. WT, wild-type.

examined these separately, viewing meal size as an index of termination and intermeal interval as an index of meal initiation. The repeated cycles of eating, terminated by a particular meal size, not eating and initiating a new meal (indexed by intermeal interval), formed the backbone of our model. We assumed that meal size and meal termination were dependent upon how hungry an animal was at the time of the meal, which will be determined by how long it was since they last ate a meal. Conversely, how long an animal waits to eat will be determined by how large their last meal was. Both cost and genotype may directly impact meal size and intermeal interval as well as exhibit interactive effects. Our model is illustrated in Fig. 6A and the intent is to discern where genotype exhibits its greatest effects. Because there is a periodicity to meals, resulting in a bimodal distribution of short (during inactive period) and long (during active period) meals (Fig. 6B), time had to be incorporated into the model. To accomplish this, we included covariance structures in the model (see Materials and methods), accounting for covariance arising from (i) individual mouse subjects (M matrix), (ii) meal start time (T matrix) and (iii) the interaction between mouse and start time (M*T).

Figures 7 and 8 show the model fits for meal size and intermeal interval, respectively, with columns showing actual data, predicted values and residuals, respectively, plotted against cost (top) and time of day (bottom). On visual inspection, the model captured the distribution of the data quite well. Moreover, the model recapitulated both the temporal and cost-related patterns of the data. Examination of residuals against several variables (e.g. genotype, cost, time of day) indicated that the variance between actual and predicted values was evenly distributed (data not shown). To assess the goodness of fit of the total model and the necessity of each factor included, we evaluated a series of models in which parameters were successively subtracted, and compared each of these partial models with the full model using a chi-square test of significance (see Materials and methods). In the meal size model, all factors were significant. In the intermeal model, the genotype × cost and cost × prior size interactions did not contribute

significantly to the model. Consequently, they were removed (resulting in no significant difference between the full and modified model with two less parameters) and the statistics reported below were derived from the modified model.

Increased dopamine delays termination of meals

The regressor coefficients and P-values associated with the different factors and their interactions are presented in Table 2. Examining first the model of meal size (termination), the preceding intermeal duration was a significant predictor of subsequent meal size but in the opposite direction from what was predicted, i.e. longer intermeal durations were associated with shorter meals. This inverse relationship between intermeal duration and meal size may reflect an underlying energy balance regulation (see below) such that, as costs increase, the subject reduces consumption and associated energy expenditure, observed in the cost main effect and delays in initiating new meals. The positive coefficient for the interaction between intermeal duration and cost suggested that as both cost and time since the last meal increased, the degree to which these two factors predicted a smaller meal was diminished, perhaps reflecting accumulating energetic urgency. Genotype exerted a significant main effect on meal size, with wildtype mice exhibiting reduced meal size, consistent with the observations above. In addition, the significant $cost \times genotype$ interaction indicated that wild-type mice showed greater declines in meal size in response to cost, consistent with the greater cost sensitivity observed in Fig. 5B. Finally, wild-type mice exhibited a significant positive offset to the negative main effect of intermeal duration [i.e. intermeal duration × genotype (wild-type)]. The interpretation of this is unclear.

In contrast to the meal size/termination model, in the model of intermeal duration (meal initiation), cost was clearly the primary factor controlling initiation, consistent with an underlying energy conserva-

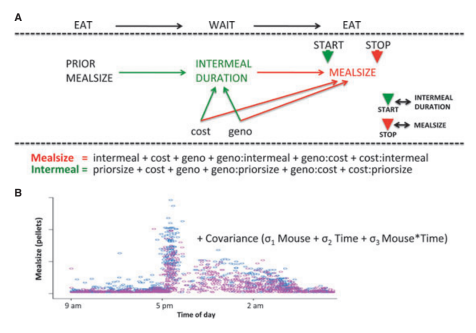


FIG. 6. Model schematic. (A) Starting and stopping meals were modeled separately with meal start (green arrow) indexed by intermeal duration and meal end (red arrow) indexed by meal size. Factors hypothesized to contribute to the start and stop models are indicated in green and red, respectively. Fixed effects are identified under the graphic. (B) A dot plot showing distribution of meal size for all mice across the experiment as a function of time of day. To capture this time-dependent distribution of meal size, covariance was explicitly modeled as indicated.

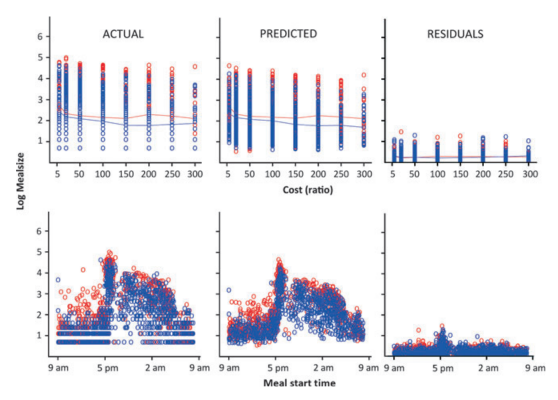


FIG. 7. Performance of meal termination (meal size) model. Actual data, predicted values and residuals presented left to right. Each point represents a meal within the entire dataset (all subjects, all days). Top row plotted by cost, bottom row plotted by meal start time. Wild-type (red) and DATkd (blue) mice. N = 8.

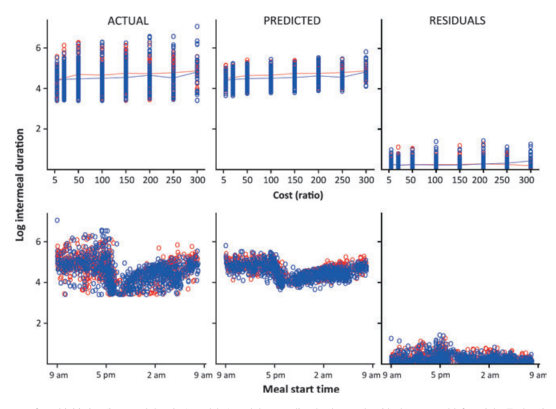


FIG. 8. Performance of meal initiation (intermeal duration) model. Actual data, predicted values and residuals presented left to right. Each point represents a meal within the entire dataset (all subjects, all days). Top row plotted by cost, bottom row plotted by meal start time. Wild-type (red) and DATkd (blue) mice. N = 8.

tion process. The lack of a significant genotype \times cost interaction was consistent with the hypothesis that both genotypes were equally subject to this conservation process. Surprisingly, prior meal size was

not a significant predictor of when a subject will initiate a new meal although we observed a significant interaction term between prior meal size and genotype. We interpret this interaction as the indirect

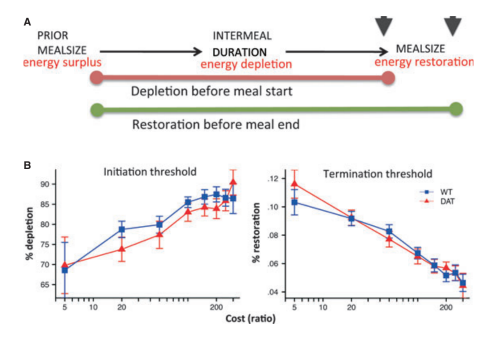


FIG. 9. Energy balance dynamics. (A) Schematic of indices used to characterize energy balance dynamics (see Materials and methods). Meal initiation indicated by green arrow, meal termination by red arrow. (B) Percent depletion of prior energy stores before initiating a new meal (left) and percent restoration of energy stores prior to terminating a meal (right). No significant differences between genotypes. N = 8. WT, wild-type.

TABLE 2. Regressor coefficients and *P*-value for models of meal size (meal termination) and intermeal duration (meal initiation)

Model	Factor	Coefficient	P-value
y = meal size	Intermeal duration	-0.0050	< 0.0001
	Cost	-0.0020	< 0.0001
	Genotype (WT)	-0.2987	0.0043
	$Cost \times genotype (WT)$	-0.0021	< 0.0001
	Intermeal duration × genotype (WT)	0.0011	0.0169
	Intermeal duration × cost	0.000015	< 0.0001
y = intermeal	Prior meal size	0.0004	0.2873
duration	Cost	0.0011	< 0.0001
	Genotype (WT)	-0.0175	0.3940
	Prior meal size – genotype (WT)	-0.0045	0.0379

WT, wild-type.

inclusion in this model of the genotype main effect observed in the meal size model.

A clear pattern emerged where cost exerted strong, highly significant main effects on both the initiation (intermeal duration) and termination (meal size) of meals, consistent with increased energy conservation as cost escalated. Genotype, however, appeared to exert its influence only on meal size/termination, including a main effect and interactions with other factors. This divergence in factors controlling the initiation and termination of meals suggested two potentially independent processes, with the termination process being dopamine dependent, whereas the initiation process appeared dopamine independent. One possibility is that the dopamine-dependent process reflected engagement of the incentive system in the pursuit of a goal, i.e. once a meal was initiated. Hyperdopaminergia may therefore facilitate goal pursuit resulting in larger meals. However, the

present data suggested that this did not significantly shift the energy balance, including conservation in the face of scarcity. Episodes of goal pursuit, i.e. meals, occurred within an overarching homeostatic system that compensated for larger meals with delayed initiation of the next meal. To assess this notion, we compared the two genotypes on energy balance dynamics.

Increased dopamine and altered meal patterning does not alter the gross rate of energy depletion or restoration

We used two indexes to assess the rate of energy depletion and restoration (Fig. 9A). First, energy depletion was calculated for the beginning of each meal as the size of the prior meal divided by the intermeal interval, approximating the degree to which an energy surplus was depleted before initiating a new meal. Second, energy restoration was calculated at the end of each meal as the size of the current meal divided by the total time since the end of the last meal (sum of preceding intermeal interval and current meal duration), i.e. the degree to which new consumption compensated for accumulating energy depletion since the last meal and restored a previous surplus.

As cost increased, both groups delayed meal initiation until greater levels of energy depletion were reached (Fig. 9B, left; cost main effect, t=4.66, P<0.0001) and terminated meals at lower levels of restoration (Fig. 9B, right; cost main effect, t=8.19, P<0.0001), reflecting a clear shift toward greater energy conservation and a net decrease in energy balance, as observed in Fig. 3. No genotype differences were observed in either of these measures (Fig. 9B, genotype main effect, depletion, t=1.18, P=0.2571; restoration, t=0.111, P=0.9126). Viewing the meal data from the perspective of decreasing and increasing energy stores, the DATkd mice showed identical patterns of energy depletion before initiating a meal as well as identical patterns of energy restoration before terminating a meal. These data suggested a level of homeostatic regulation of energy balance that was unaffected by elevated dopamine. Although

increased dopamine delayed meal termination (prolonged pursuit) and changed meal patterning, energy balance dynamics were identical between the groups. This explains why, despite increased within-meal motivation/effort, increased dopamine did not substantially impact energy balance and survival in response to escalating cost; both the DATkd and wild-type mice were equally subject to homeostatic conservation mechanisms induced by increasing caloric restriction and food scarcity in the demand paradigm used here.

Increased dopamine enhances effort in pursuit of individual meals without altering overall consumption

To address the role of dopamine in meal patterning without the confound of increasing scarcity and putative homeostatic conservation mechanisms, we used an alternative homecage progressive ratio paradigm in which increasing costs were incorporated into individual meals. In this way, each meal indicated a subject's willingness to continue working for food. However, after stopping a meal, the cost reset, allowing the animal access once again to lower cost food. In this way, mice could maintain their energy balance by eating more frequent, smaller meals, eliminating caloric restriction and implicit food scarcity. In the course of the study, both groups reduced their initial body weight by 7-8% with no differences between genotypes (t = 0.427, P = 0.6801, data not shown). There was, however, a significant difference between the groups in initial weight (means, DAT = 25.7 g, wild-type = 21.4 g, P = 0.008). Consequently, we used two-way ANOVA with initial body weight and genotype as factors, except for the number of meals where only genotype was included. No significant interaction effects were observed. The P-values in Fig. 10 indicate genotype main effect.

In this paradigm, the cost of a 20 mg pellet incremented by 2 after each pellet earned (see Materials and methods). The incrementing schedule reset after 30 min of inactivity. Highly motivated mice may eat large meals. Less motivated mice may eat smaller meals. However, by eating a greater number of small meals, mice can easily maintain their desired intake. Consistent with the meal patterning results above, the DATkd mice showed a statistically significant increase in their breakpoint (Fig. 10A; $F_{1,8} = 7.86$, P = 0.0263), meal size and meal duration (Fig. 10B; meal size, $F_{1,8} = 6.02$, P = 0.0438; duration, $F_{1,8} = 6.74$, P = 0.0355), with a corresponding marginally significant decrease in their total number of meals (Fig. 10B, $F_{1,8} = 4.19$, P = 0.0709). However, their overall consumption was essentially equivalent (Fig. 10A; $F_{1.8} = 0.988$, P = 0.3533), although they worked considerably more to maintain the same energy balance (Fig. 10A; $F_{1,8} = 16.9$, P = 0.0045). These data were consistent with the above results where increased dopamine affected motivation and willingness to work within individual meals (episodes of goal pursuit) presumably without altering the homeostatically regulated overall energy balance.

Finally, to discern whether the 'cost' associated with pressing reflected the effort required or the delay associated with obtaining a reward as the number of required lever presses increased, we tested a homecage progressive interval paradigm where a single press always resulted in a pellet, but an incrementing delay was introduced between pellets (incrementing by 30 s). In this paradigm, no differences were observed between genotypes (Fig. 11), indicating that within these homecage studies it was effort rather than delay that comprised cost (Floresco et al., 2008). In this study, initial weights were identical between groups (t = 0.0511, P = 0.9603).

Discussion

Dopamine is thought by many to enhance incentive motivation and willingness to work toward a goal, an idea that has linked it to addiction and compulsive drug-seeking (Berridge, 2007; Di Chiara & Bassareo, 2007) and, more recently, the regulation of food intake and obesity (Volkow & Wise, 2005; Kenny, 2010; Volkow et al., 2010). In the DATkd mice with elevated tonic dopamine, we did observe an increase in responding at higher costs, but this increase over wild-type mouse responding was small compared with the decline observed in

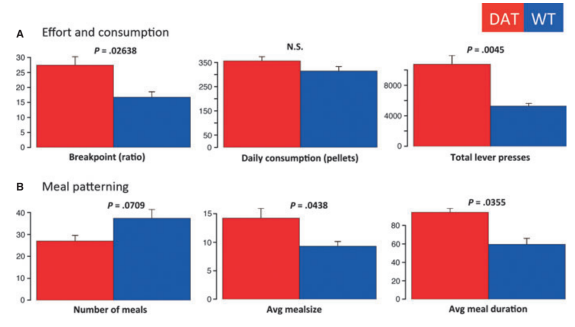


FIG. 10. Homecage progressive ratio. (A) Comparison of genotypes on breakpoint (left, defined as last ratio completed before pressing stopped for 30 min and incrementing ratio reset), total daily consumption (middle) and total daily lever presses (right). (B) Comparison of genotype on average number of meals (left), average meal size (middle) and average meal duration (right). N = 5. WT, wild-type.

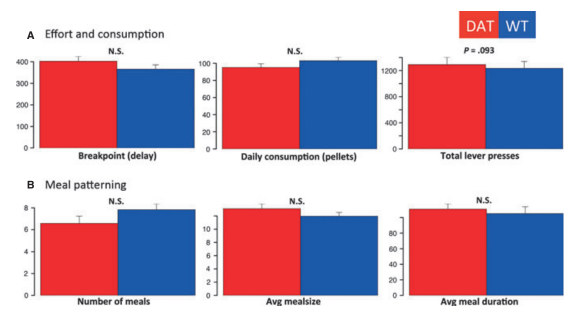


FIG. 11. Homecage progressive interval. (A) Comparison of genotypes on breakpoint (left, defined as last interval before pressing stopped for 30 min and incrementing interval reset), total daily consumption (middle) and total daily lever presses (right). (B) Comparison of genotype on average number of meals (left), average meal size (middle) and average meal duration (right). N = 5. WT, wild-type.

both genotypes as costs escalate. Critically, this difference did not alter the adaptive survival and comparison of elasticity between groups, using an established model of demand elasticity, showed no significant genotype difference. These data, together with those showing no difference between genotypes at baseline, low-cost conditions, suggest that dopaminergic modulation of incentive and willingness to work is subject to other processes regulating food intake, presumably homeostatic mechanisms. In the demand study, we suggest that increasing caloric restriction resulting from an escalating work requirement induces food scarcity and engages mechanisms of homeostatic energy conservation, although elaborating such mechanisms is beyond the scope of the present study.

Although we cannot conclusively rule out in the demand study that the energy required to obtain pellets at higher ratios exceeds the energy obtained, this is unlikely for several reasons. First, those mice that persevered at higher ratios were able to maintain a reduced body weight, suggesting that those that dropped out at lower costs were not simply responding to an energetic deficit. Second, the homecage progressive ratio shows the same pattern of results (different meal patterning between the genotypes with similar overall energy balance) without the problem of a potential energetic ceiling. Finally, even if the demand curve partially reflects diminishing caloric gain from each pellet earned, the mice still show a differential response to this diminishing return that reflects their motivation to persevere in the face of those costs and the implicit scarcity that they induce.

In contrast to the overall energy balance, in which the two genotypes are more similar than different across costs, hyperdopaminergia has a clear effect on meal patterning with DATkd mice eating larger but fewer meals. This effect is more pronounced as the cost of food increases, with the DATkd mice showing relative insensitivity to cost in terms of meal size, consistent with views that dopamine increases incentive and willingness to work. However, this effect does not appear to significantly alter the energy balance as prolonged intermeal intervals compensate for larger meals, resulting in a similar net caloric intake. Modeling of the meal data indicates that significant genotype effects center upon delaying the termination of a meal

(i.e. increasing meal size) with no direct effects on meal initiation (i.e. intermeal interval). Whether dopamine is acting to enhance a positive incentive value or diminish the impact of increasing satiety was not specifically addressed by the present study. However, the lack of difference between genotypes in the homecage progressive interval suggests that elevated dopamine is not acting to diminish satiety signals. If this were the case, we might also expect to see prolonged meals in the progressive interval study.

Taken as a whole, these data suggest that increased dopamine can clearly and significantly increase incentive and willingness to work but that this effect is constrained to temporally local episodes of goal pursuit, in this case a meal [see Desrochers et al. (2010) for a related study of the localization of sensitivity to costs in reinforcement learning]. Put another way, dopamine does not appear to globally enhance incentive value and propensity to work but rather modulates local effort once goal pursuit is initiated ('in the heat of the meal'), leaving the overall homeostatic control of energy balance intact. As a consequence, hyperdopaminergia prolongs meals, thus increasing meal size. The net energy balance is maintained, however, with a decrease in meal frequency, suggesting that dopamine is not playing a significant role in initiating meals. Interestingly, Zorrilla et al. (2005) found the opposite pattern where the administration of acute leptin primarily reduced the frequency but not size of meals. Importantly, these conclusions may not apply across the entire range of dopamine concentrations. For example, a complete lack of dopamine in the dorsal striatum eliminates feeding behavior altogether (Palmiter, 2008). At lower dopamine concentrations, meal initiation may be altered, although whether the frequency would be decreased [as suggested by the work of both Palmiter (2008) and Salamone et al. (1990, 1991)] or increased [as proposed by the 'reward deficiency' hypothesis (Kenny, 2010)] will require further investigation. At lower dopamine concentrations, dopamine's role in motor control may also impact the initiation of meals, i.e. 'meal initiating akinesia'.

Several limitations need to be noted. First, although the present findings are consistent with the role ascribed to the striatum in appetitive motivation, we cannot attribute the present observations to a

discrete striatal region. Both the dorsal and ventral striatum have been implicated in control of feeding (Kelley, 2004; Balleine et al., 2007; Palmiter, 2008) and both are affected in the DATkd mice. In addition, although the effects are less striking than in the striatum, the DATkd mice show increased dopamine turnover in the hypothalamus (see Materials and methods), leaving open the possibility that hypothalamic dopamine contributes to the present observations. Interestingly, if so, this would suggest that hypothalamic dopamine is not contributing to the overall homeostatic regulation of energy balance. A more likely interpretation is that the alterations in hypothalamic dopamine are marginal and have little impact on behavior and the dramatic changes in striatal dopamine account for the observed changes in appetitive and instrumental responding, consistent with decades of literature on the role of striatum in mediating these behaviors. Importantly, a critical component of the addiction perspective is the loss of prefrontal cortex executive control inhibiting compulsive responding (George & Koob, 2010; Koob & Volkow, 2010). In the DATkd mice, altered dopamine function in the prefrontal cortex (PFC) is unlikely to contribute to the present observations as dopamine reuptake in the PFC is mediated primarily by the norepinephrine transporter and alterations in DAT have little impact on PFC dopamine reuptake kinetics (Sesack et al., 1998; Mundorf et al., 2001; Moron et al., 2002). Thus, the increased meal size observed here is unlikely to be attributable to a loss of inhibitory cortical control but rather reflects enhanced incentive via striatal signals or, possibly, hypothalamic signals.

Second, we examined hyperdopaminergia but not hypodopaminergia and cannot assume a simple inversion of the results. Although the work of Salamone et al. (1990, 1991) demonstrates clearly that decreased dopamine can diminish effort, the effect of decreased dopamine on meal patterns in a semi-naturalistic environment remains to be tested empirically. The present data, nonetheless, are difficult to reconcile with the 'dopamine deficiency' hypothesis currently gaining prominence (Kenny, 2010). In this theory, reduced dopamine function causes a deficiency in reward signaling, leading to increased consumption in order to compensate for this reduction in reward. If dopamine signaling provided direct reward experience as suggested by this hypothesis, we would expect that hyperdopaminergic mice would decrease their consumption due to an analogous 'dopamine excess,' which they do not.

Third, we used a standard grain diet. It is widely proposed that palatable, energy-rich foods are highly rewarding and promote consumption beyond homeostatically regulated energy needs (Saper et al., 2002; Zheng et al., 2009; Kenny, 2010; Oswald et al., 2010; Volkow et al., 2010). Under these circumstances, it may be that hyperdopaminergia would not only increase meal size but also foodseeking and meal initiation, significantly contributing to a positive energy balance and homeostatic dysregulation. Finally, in the mice tested here, there is no known pathophysiology in neuroendocrine, homeostatic systems, such as deficits in leptin signaling. It is possible that, in the context of dysregulated homeostatic mechanisms, dopamine may exert different, unexpected effects on both energy balance and meal patterning. Addressing these issues requires further study.

We propose the following simplified model. As energy stores are depleted, hunger signals increase until reaching a threshold that initiates food-seeking and consumption. As energy is ingested, metabolic signals indicate rising energy levels, moving the organism back below the meal initiation threshold. If only a single threshold controlled consumption, then, shortly after initiating a meal, a signaled rise in energy (e.g. blood sugar, insulin) would drop below the 'initiating threshold' and the animal would discontinue eating after only a short period of ingestion. A single energy threshold for eating/not eating would result in distributed, constant grazing, precluding the ability to store energy, decreasing the animal's ability to exploit found food sources and increasing the amount of time spent foraging, notably increasing the risk of predation. However, establishing two energy thresholds, one for initiating and one for terminating a meal, would alleviate this problem. Extracellular dopamine increases after the initiation of a meal and remains elevated for 20-60 min (Hernandez & Hoebel, 1988; Hoebel et al., 1992; Yoshida et al., 1992; Wilson et al., 1995; Martel & Fantino, 1996; Taber & Fibiger, 1997; Sokolowski et al., 1998; Cousins et al., 1999; Ostlund et al., 2010). We suggest that this increase in dopaminergic tone after initiating a meal enhances the incentive associated with pursuit and consumption, effectively generating a different 'termination' threshold, prolonging motivation despite signals indicating increasing energy stores, enabling the animal to consume a larger meal. This would result in larger but fewer meals. The resulting intermeal interval could be used for other activities, including further exploration. The lack of a genotype difference in the homecage progressive interval study suggests that this dopaminergic setting of a 'termination' threshold may arise through enhancing willingness to expend energy in pursuit of a goal rather than through directly diminishing the effects of satiety.

Although making inferences from a mouse study to processes contributing to human behavior and obesity requires caution, the present study suggests a couple of potentially important points.

The present data suggest that increased reward/dopaminergic function would prolong individual episodes of eating (meals/snacks), increasing caloric intake, without necessarily increasing the frequency of eating. Human eating, however, is not entirely homeostatic (Strubbe & Woods, 2004; Levitsky, 2005; Lowe & Levine, 2005; Woods & D'Alessio, 2008; Grill, 2010), i.e. we do not eat only when hungry, but at scheduled meal times, in social settings when food is provided (e.g. donuts in the office, celebrations) and for a variety of other reason, including stress and anxiety (Dallman, 2010). Moreover, caloric consumption is not limited to meals but often occurs almost continuously throughout the day in the form of calorie-rich drinks such as soft drinks and lattes. Under these circumstances, enhanced dopamine function may prolong ingestion during any single episode of consumption (the 'one more potato chip' phenomenon) without a corresponding increase in wait time prior to the next meal or snack, resulting in a net positive energy balance. Individual differences in dopamine function and reward sensitivity may contribute to vulnerability to overeating and obesity (Campbell & Eisenberg, 2007) by enhancing prolongation of ingestion. In addition, as frequently suggested, the prevalence of and easy access to highly palatable, energy-rich foods may itself contribute to enhanced dopaminergic responses to food-related stimuli (Hajnal & Norgren, 2002; Small et al., 2003; Avena et al., 2008; Davis & Carter, 2009) generating dopaminergic dysregulation. The present data do not, however, suggest that elevated dopamine would increase food-seeking although, as noted earlier, whether this holds true with high-sugar and high-fat food remains to be tested.

In contrast, as weight accumulates and overweight transitions into obesity, an opposite mechanism may be engaged. In this case, elevated leptin may result in two physiological adaptations: (i) leptin receptor insensitivity (Considine et al., 1996; Maffei et al., 1996; Munzberg et al., 2004; Farooqi & O'Rahilly, 2005; Enriori et al., 2007; Myers et al., 2008; Opland et al., 2010) and (ii) diminished dopamine function (Fulton et al., 2006; Roseberry et al., 2007). Under these conditions, prolongation of consumption during individual episodes of ingestion might be replaced by greater food-seeking and increased frequency of consumption as the homeostatic system behaves as if it were in energy deficit (i.e. reduced leptin signaling), keeping the goal of energy acquisition consistently active. Diminished dopaminergic function may result in diminished motivation to overcome costs in these individuals; however, cost is rarely an issue in our modern culture where energy-dense foods are cheap, ubiquitous and plentiful. This view would suggest that these two populations (overweight and obese) may benefit from different interventions, both behaviorally and pharmacologically. To our knowledge, there have been no human studies that investigate meal patterning across the natural course of obesity, i.e. from overweight to obese. The present study suggests that dopaminergic effects on meal patterning could play a role in the transition from overweight to obese and suggest a specific way in which dopaminergic, incentive mechanisms and environmental factors may interact to promote net positive energy balance, weight gain and obesity.

Supporting Information

Additional supporting information can be found in the online version of this article:

Fig. S1. Plots of consumption by cost for individual mice. WT, wild-type; 'rew', reward.

Fig. S2. Activity patterns. (A) Average daily inactive lever presses (LP) at each ratio/cost. (B) Average daily within-meal lever-pressing rate. (C) Histogram of inter-response times (IRTs) as percentage of total IRTs in 1 s bins. (D) Histogram of post-reinforcement pauses (PRPs) in 10 s bins from 1 to 600 s (5 min). No significant genotype main effect or interactions observed. N = 8. WT, wild-type.

Please note: As a service to our authors and readers, this journal provides supporting information supplied by the authors. Such materials are peer-reviewed and may be re-organized for online delivery, but are not copy-edited or typeset by Wiley-Blackwell. Technical support issues arising from supporting information (other than missing files) should be addressed to the authors.

Acknowledgements

This work was supported by NIDA (DA25875 to J.A.B.), NIDDK (R56DK088515 to X.Z.), NARSAD (to X.Z.) and NIDA (F31DA026802 to C.R.M.F.). We would like to thank the University of Chicago Department of Statistics consulting program, particularly Peter McCullagh, Duo Jiang and Mei Wang for their assistance with modeling the meal data.

Abbreviations

DAT, dopamine transporter; DATkd, dopamine transporter knock-down; FR, fixed-ratio.

References

- Avena, N.M., Rada, P. & Hoebel, B.G. (2008) Evidence for sugar addiction: behavioral and neurochemical effects of intermittent, excessive sugar intake. *Neurosci. Biobehav. Rev.*, **32**, 20–39.
- Balleine, B.W. (2005) Neural bases of food-seeking: affect, arousal and reward in corticostriatolimbic circuits. *Physiol. Behav.*, **86**, 717–730.
- Balleine, B.W., Delgado, M.R. & Hikosaka, O. (2007) The role of the dorsal striatum in reward and decision-making. *J. Neurosci.*, **27**, 8161–8165.
- Beeler, J.A., Daw, N., Frazier, C.R. & Zhuang, X. (2010) Tonic dopamine modulates exploitation of reward learning. *Front. Behav. Neurosci.*, 4, 170.
- Berridge, K.C. (2007) The debate over dopamine's role in reward: the case for incentive salience. *Psychopharmacology (Berl.)*, **191**, 391–431.
- Berridge, K.C., Ho, C.Y., Richard, J.M. & DiFeliceantonio, A.G. (2010) The tempted brain eats: pleasure and desire circuits in obesity and eating disorders. *Brain Res.*, **1350**, 43–64.
- Berthoud, H.R. (2007) Interactions between the "cognitive" and "metabolic" brain in the control of food intake. *Physiol. Behav.*, **91**, 486–498.

- Berthoud, H.R. & Morrison, C. (2008) The brain, appetite, and obesity. *Annu. Rev. Psychol.*, **59**, 55–92.
- Bosse, R., Fumagalli, F., Jaber, M., Giros, B., Gainetdinov, R.R., Wetsel, W.C., Missale, C. & Caron, M.G. (1997) Anterior pituitary hypoplasia and dwarfism in mice lacking the dopamine transporter. *Neuron*, 19, 127–138
- Cagniard, B., Balsam, P.D., Brunner, D. & Zhuang, X. (2006a) Mice with chronically elevated dopamine exhibit enhanced motivation, but not learning, for a food reward. *Neuropsychopharmacology*, 31, 1362–1370.
- Cagniard, B., Beeler, J.A., Britt, J.P., McGehee, D.S., Marinelli, M. & Zhuang, X. (2006b) Dopamine scales performance in the absence of new learning. Neuron, 51, 541–547.
- Campbell, B.C. & Eisenberg, D. (2007) Obesity, attention deficit-hyperactivity disorder and the dopaminergic reward system. Coll. Antropol., 31, 33–38.
- Center for Disease Control and Prevention. (2010) *U.S. Obesity Trends: Trends by State 1985–2010*. Available at http://www.cdc.gov/obesity/data/trends. html.
- Chaney, M.A. & Rowland, N.E. (2008) Food demand functions in mice. *Appetite*, **51**, 669–675.
- Considine, R.V., Sinha, M.K., Heiman, M.L., Kriauciunas, A., Stephens, T.W., Nyce, M.R., Ohannesian, J.P., Marco, C.C., McKee, L.J., Bauer, T.L. & Caro, J.F. (1996) Serum immunoreactive-leptin concentrations in normal-weight and obese humans. N. Engl. J. Med., 334, 292–295.
- Cousins, M.S., Trevitt, J., Atherton, A. & Salamone, J.D. (1999) Different behavioral functions of dopamine in the nucleus accumbens and ventrolateral striatum: a microdialysis and behavioral investigation. *Neuroscience*, 91, 925–934.
- Dallman, M.F. (2010) Stress-induced obesity and the emotional nervous system. *Trends Endocrinol. Metab.*, 21, 159–165.
- Davis, C. & Carter, J.C. (2009) Compulsive overeating as an addiction disorder. A review of theory and evidence. Appetite, 53, 1–8.
- Davis, J.F., Choi, D.L., Schurdak, J.D., Fitzgerald, M.F., Clegg, D.J., Lipton, J.W., Figlewicz, D.P. & Benoit, S.C. (2010) Leptin regulates energy balance and motivation through action at distinct neural circuits. *Biol. Psychiatry*, 69, 668–674
- Dayan, P. & Balleine, B.W. (2002) Reward, motivation, and reinforcement learning. Neuron, 36, 285–298.
- Desrochers, T.M., Jin, D.Z., Goodman, N.D. & Graybiel, A.M. (2010) Optimal habits can develop spontaneously through sensitivity to local cost. *Proc. Natl. Acad. Sci. USA*, **107**, 20512–20517.
- Di Chiara, G. (2005) Dopamine in disturbances of food and drug motivated behavior: a case of homology? *Physiol. Behav.*, **86**, 9–10.
- Di Chiara, G. & Bassareo, V. (2007) Reward system and addiction: what dopamine does and doesn't do. Curr. Opin. Pharmacol., 7, 69–76.
- Enriori, P.J., Evans, A.E., Sinnayah, P., Jobst, E.E., Tonelli-Lemos, L., Billes, S.K., Glavas, M.M., Grayson, B.E., Perello, M., Nillni, E.A., Grove, K.L. & Cowley, M.A. (2007) Diet-induced obesity causes severe but reversible leptin resistance in arcuate melanocortin neurons. *Cell Metab.*, 5, 181–194.
- Everitt, B.J. & Robbins, T.W. (2005) Neural systems of reinforcement for drug addiction: from actions to habits to compulsion. *Nat. Neurosci.*, **8**, 1481–1489
- Farooqi, I.S. & O'Rahilly, S. (2005) Monogenic obesity in humans. *Annu. Rev. Med.*, **56**, 443–458.
- Figlewicz, D.P. & Sipols, A.J. (2010) Energy regulatory signals and food reward. *Pharmacol. Biochem. Behav.*, **97**, 15–24.
- Figlewicz, D.P., MacDonald Naleid, A. & Sipols, A.J. (2007) Modulation of food reward by adiposity signals. *Physiol. Behav.*, **91**, 473–478.
- Floresco, S.B., Tse, M.T. & Ghods-Sharifi, S. (2008) Dopaminergic and glutamatergic regulation of effort- and delay-based decision making. *Neuropsychopharmacology*, **33**, 1966–1979.
- Fulton, S., Pissios, P., Manchon, R.P., Stiles, L., Frank, L., Pothos, E.N., Maratos-Flier, E. & Flier, J.S. (2006) Leptin regulation of the mesoaccumbens dopamine pathway. *Neuron*, 51, 811–822.
- George, O. & Koob, G.F. (2010) Individual differences in prefrontal cortex function and the transition from drug use to drug dependence. *Neurosci. Biobehav. Rev.*, **35**, 232–247.
- Grill, H.J. (2010) Leptin and the systems neuroscience of meal size control. Front. Neuroendocrinol., 31, 61–78.
- Hajnal, A. & Norgren, R. (2002) Repeated access to sucrose augments dopamine turnover in the nucleus accumbens. *Neuroreport*, 13, 2213–2216.
- Hernandez, L. & Hoebel, B.G. (1988) Feeding and hypothalamic stimulation increase dopamine turnover in the accumbens. *Physiol. Behav.*, 44, 599–606.
- Hoebel, B.G., Mark, G.P. & West, H.L. (1992) Conditioned release of neurotransmitters as measured by microdialysis. *Clin. Neuropharmacol.*, 15(Suppl 1 Pt A), 704A–705A.

- Hommel, J.D., Trinko, R., Sears, R.M., Georgescu, D., Liu, Z.W., Gao, X.B., Thurmon, J.J., Marinelli, M. & DiLeone, R.J. (2006) Leptin receptor signaling in midbrain dopamine neurons regulates feeding. Neuron, 51, 801-810.
- Hursh, S.R. & Silberberg, A. (2008) Economic demand and essential value. Psychol. Rev., 115, 186–198.
- Kelley, A.E. (2004) Ventral striatal control of appetitive motivation: role in ingestive behavior and reward-related learning. Neurosci. Biobehav. Rev., **27**. 765–776.
- Kelley, A.E., Baldo, B.A. & Pratt, W.E. (2005a) A proposed hypothalamicthalamic-striatal axis for the integration of energy balance, arousal, and food reward. J. Comp. Neurol., 493, 72-85.
- Kelley, A.E., Baldo, B.A., Pratt, W.E. & Will, M.J. (2005b) Corticostriatalhypothalamic circuitry and food motivation: integration of energy, action and reward. Physiol. Behav., 86, 773-795.
- Kenny, P.J. (2010) Reward mechanisms in obesity: new insights and future directions. Neuron, 69, 664-679.
- Koob, G.F. & Volkow, N.D. (2010) Neurocircuitry of addiction. Neuropsychopharmacology, 35, 217-238.
- Levitsky, D.A. (2005) The non-regulation of food intake in humans: hope for reversing the epidemic of obesity. Physiol. Behav., 86, 623-632.
- Lowe, M.R. & Levine, A.S. (2005) Eating motives and the controversy over dieting: eating less than needed versus less than wanted. Obes. Res., 13, 797-
- Lutter, M. & Nestler, E.J. (2009) Homeostatic and hedonic signals interact in the regulation of food intake. J. Nutr., 139, 629-632.
- Maffei, M., Stoffel, M., Barone, M., Moon, B., Dammerman, M., Ravussin, E., Bogardus, C., Ludwig, D.S., Flier, J.S., Talley, M., Auerbach, S. & Friedman, J.M. (1996) Absence of mutations in the human OB gene in obese/diabetic subjects. Diabetes, 45, 679-682.
- Martel, P. & Fantino, M. (1996) Mesolimbic dopaminergic system activity as a function of food reward: a microdialysis study. Pharmacol. Biochem. Behav., 53, 221-226.
- Moron, J.A., Brockington, A., Wise, R.A., Rocha, B.A. & Hope, B.T. (2002) Dopamine uptake through the norepinephrine transporter in brain regions with low levels of the dopamine transporter: evidence from knock-out mouse lines. J. Neurosci., 22, 389-395.
- Mundorf, M.L., Joseph, J.D., Austin, C.M., Caron, M.G. & Wightman, R.M. (2001) Catecholamine release and uptake in the mouse prefrontal cortex. J. Neurochem., 79, 130-142.
- Munzberg, H., Flier, J.S. & Bjorbaek, C. (2004) Region-specific leptin resistance within the hypothalamus of diet-induced obese mice. Endocrinology, 145, 4880-4889.
- Myers, M.G., Cowley, M.A. & Munzberg, H. (2008) Mechanisms of leptin action and leptin resistance. Annu. Rev. Physiol., 70, 537-556.
- Neel, J.V. (1999) The "thrifty genotype" in 1998. Nutr. Rev., 57, S2-S9.
- Opland, D.M., Leinninger, G.M. & Myers, M.G. Jr (2010) Modulation of the mesolimbic dopamine system by leptin. Brain Res., 1350, 65-70.
- Ostlund, S.B., Wassum, K.M., Murphy, N.P., Balleine, B.W. & Maidment, N.T. (2010) Extracellular dopamine levels in striatal subregions track shifts in motivation and response cost during instrumental conditioning. J. Neurosci., 31, 200-207.
- Oswald, K.D., Murdaugh, D.L., King, V.L. & Boggiano, M.M. (2010) Motivation for palatable food despite consequences in an animal model of binge eating. Int. J. Eat. Disord., 44, 203-211.
- Palmiter, R.D. (2007) Is dopamine a physiologically relevant mediator of feeding behavior? Trends Neurosci., 30, 375-381.
- Palmiter, R.D. (2008) Dopamine signaling in the dorsal striatum is essential for motivated behaviors: lessons from dopamine-deficient mice. Ann. NY Acad. Sci., 1129, 35-46.
- Pecina, S., Cagniard, B., Berridge, K.C., Aldridge, J.W. & Zhuang, X. (2003) Hyperdopaminergic mutant mice have higher "wanting" but not "liking" for sweet rewards. J. Neurosci., 23, 9395-9402.
- Redish, A.D. (2004) Addiction as a computational process gone awry. Science, 306, 1944-1947.
- Roseberry, A.G., Painter, T., Mark, G.P. & Williams, J.T. (2007) Decreased vesicular somatodendritic dopamine stores in leptin-deficient mice. J. Neurosci., 27, 7021-7027.
- Rowland, N.E., Vaughan, C.H., Mathes, C.M. & Mitra, A. (2008) Feeding behavior, obesity, and neuroeconomics. Physiol. Behav., 93, 97-109.

- Salamone, J.D. (2007) Functions of mesolimbic dopamine: changing concepts and shifting paradigms. Psychopharmacology (Berl.), 191, 389.
- Salamone, J.D. & Correa, M. (2002) Motivational views of reinforcement: implications for understanding the behavioral functions of nucleus accumbens dopamine. Behav. Brain Res., 137, 3-25.
- Salamone, J.D., Zigmond, M.J. & Stricker, E.M. (1990) Characterization of the impaired feeding behavior in rats given haloperidol or dopamine-depleting brain lesions. Neuroscience, 39, 17-24.
- Salamone, J.D., Steinpreis, R.E., McCullough, L.D., Smith, P., Grebel, D. & Mahan, K. (1991) Haloperidol and nucleus accumbens dopamine depletion suppress lever pressing for food but increase free food consumption in a novel food choice procedure. Psychopharmacology (Berl.), 104, 515-521.
- Salamone, J.D., Cousins, M.S., McCullough, L.D., Carriero, D.L. & Berkowitz, R.J. (1994) Nucleus accumbens dopamine release increases during instrumental lever pressing for food but not free food consumption. Pharmacol. Biochem. Behav., 49, 25-31.
- Salamone, J.D., Cousins, M.S. & Snyder, B.J. (1997) Behavioral functions of nucleus accumbens dopamine: empirical and conceptual problems with the anhedonia hypothesis. Neurosci. Biobehav. Rev., 21, 341-359.
- Saper, C.B., Chou, T.C. & Elmquist, J.K. (2002) The need to feed: homeostatic and hedonic control of eating. Neuron, 36, 199-211.
- Schultz, W. (2007) Multiple dopamine functions at different time courses. Annu. Rev. Neurosci., 30, 259-288.
- Sesack, S.R., Hawrylak, V.A., Guido, M.A. & Levey, A.I. (1998) Cellular and subcellular localization of the dopamine transporter in rat cortex. Adv. Pharmacol., 42, 171-174.
- Small, D.M., Jones-Gotman, M. & Dagher, A. (2003) Feeding-induced dopamine release in dorsal striatum correlates with meal pleasantness ratings in healthy human volunteers. Neuroimage, 19, 1709-1715.
- Sokolowski, J.D., Conlan, A.N. & Salamone, J.D. (1998) A microdialysis study of nucleus accumbens core and shell dopamine during operant responding in the rat. Neuroscience, 86, 1001-1009.
- Strubbe, J.H. & Woods, S.C. (2004) The timing of meals. Psychol. Rev., 111, 128 - 141.
- Taber, M.T. & Fibiger, H.C. (1997) Feeding-evoked dopamine release in the nucleus, accumbens: regulation by glutamatergic mechanisms. Neuroscience, 76, 1105-1112.
- Volkow, N.D. & Wise, R.A. (2005) How can drug addiction help us understand obesity? Nat. Neurosci., 8, 555-560.
- Volkow, N.D., Wang, G.J. & Baler, R.D. (2010) Reward, dopamine and the control of food intake: implications for obesity. Trends Cogn. Sci., 15, 37-
- Wang, Y. & Beydoun, M.A. (2007) The obesity epidemic in the United States gender, age, socioeconomic, racial/ethnic, and geographic characteristics: a systematic review and meta-regression analysis. Epidemiol. Rev., 29, 6-28.
- Wilson, C., Nomikos, G.G., Collu, M. & Fibiger, H.C. (1995) Dopaminergic correlates of motivated behavior: importance of drive. J. Neurosci., 15, 5169-5178.
- Wise, R.A. (2004) Dopamine, learning and motivation. Nat. Rev. Neurosci., 5, 483-494.
- Woods, S.C. & D'Alessio, D.A. (2008) Central control of body weight and appetite. J. Clin. Endocrinol. Metab., 93, S37-S50.
- Yin, H.H., Zhuang, X. & Balleine, B.W. (2006) Instrumental learning in hyperdopaminergic mice. Neurobiol. Learn. Mem., 85, 283-288.
- Yoshida, M., Yokoo, H., Mizoguchi, K., Kawahara, H., Tsuda, A., Nishikawa, T. & Tanaka, M. (1992) Eating and drinking cause increased dopamine release in the nucleus accumbens and ventral tegmental area in the rat: measurement by in vivo microdialysis. Neurosci. Lett., 139, 73-76.
- Zheng, H., Lenard, N.R., Shin, A.C. & Berthoud, H.R. (2009) Appetite control and energy balance regulation in the modern world: reward-driven brain overrides repletion signals. Int. J. Obes. (Lond)., 33(Suppl 2), S8-S13.
- Zhuang, X., Oosting, R.S., Jones, S.R., Gainetdinov, R.R., Miller, G.W., Caron, M.G. & Hen, R. (2001) Hyperactivity and impaired response habituation in hyperdopaminergic mice. Proc. Natl. Acad. Sci. USA, 98, 1982-1987
- Zorrilla, E.P., Inoue, K., Valdez, G.R., Tabarin, A. & Koob, G.F. (2005) Leptin and post-prandial satiety: acute central leptin more potently reduces meal frequency than meal size in the rat. Psychopharmacology (Berl.), 177, 324-