Sugar and Fat Bingeing Have Notable Differences in Addictive-like Behavior

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Introduction

Although binge-eating behavior is traditionally associated with eating disorders, it is becoming more prevalent in the United States through its emergence in a variety of clinical and nonclinical populations. Binge eating has been linked to obesity, which presently afflicts 33% of the adult U.S. population (1, 2) and may also be a predictor of body-fat gain among children (3). Binge eating is also associated with increased frequency of body weight fluctuation, depression, anxiety, and substance abuse (4-6). The presence of bingeing behavior in several different eating disorders, as well as in nonclinical populations, has made it important to study from a public-health perspective.

The Diagnostic and Statistical Manual of Mental Disorders (ed. 4) defines binge eating as a series of recurrent binge episodes in which each episode is defined as eating a larger amount of food than normal during a short period of time (usually within any 2-h period) (7). Binge-eating episodes are associated with 3 or more of the following: 1) eating until feeling uncomfortably full, 2) eating large amounts of food when not physically hungry, 3) eating more rapidly than normal, 4) eating alone because one is embarrassed by how much s/he is eating, 5) feeling disgusted, depressed, or guilty after overeating, or 6) marked distress or anxiety regarding binge eating.

Aside from diagnosed patients, there is also a far larger population of individuals who often binge on food, but perhaps not regularly enough to warrant a clinical diagnosis. It is not always clear where one draws the line between simply eating a large meal and a pathological binge. However, the physiological consequences of binge eating may be similar, whether engaged in naturally because of hunger, casually for social or hedonic reasons, or regularly enough to warrant a diagnosis.

What are the common binge foods?

To put it simply, people usually binge on highly palatable energy-rich food. These foods are typically high in fats, sugars, or often both (8, 9). Binge episodes often involve consumption of bread or pasta, followed in frequency by sweets, fatty foods, or salty snacks (10). Individuals with a preference for binging on sweet foods tend to binge more frequently.

Why do people not binge on broccoli? There must be some property of palatable “dessert” and “snack” foods rich in sugar and/or fat that promotes binge eating. Sugars and fats are well known to have different effects on physiology and brain chemistry (11), which may be related to their different effects on behavior. To understand the behavioral and neurochemical basis of binge eating on specific macronutrients, we turn to laboratory animal models of binge eating.

Animal models of binge eating

Binge eating is a multifaceted behavior, with emotional and cultural components that are difficult to reproduce with animal models. Nonetheless, animal models of binge eating are funda-
Models of sugar bingeing
Several laboratories have used limited access to sugar solutions to model binge eating (12–15). The findings all suggest that animals will engage in binge-type eating on a sweet food when it is offered intermittently. Our laboratory has developed a model of sugar bingeing (16) in which rats are maintained on daily 12-h food restriction, followed by access to a 25% glucose or 10% sucrose solution (similar to the sugar concentration of a soft drink) and a nonpurified rodent diet. After a few days on this schedule, these rats escalate their daily intake of sugar (Fig. 1 A) and begin to binge, as indicated by an increase in their intake of the sugar solution during the first hour of access. Rats that have access to the sugar solution and nonpurified diet ad libitum consume a total daily amount similar to that consumed by the bingeing rats, but they seldom engage in discrete bingeing episodes. Body weight and total daily caloric intake do not differ from normal in rats that are bingeing on sugar (Fig. 1 C), indicating that the rats are able to regulate their energy intake and compensate for the excess energy by eating less rodent nonpurified diet (Fig. 1 B).

Models of fat bingeing
Animals will also binge on pure fat, which suggests that binge eating is not exclusive to sweet taste. Corwin et al. (17) have shown that sated rats with access to rodent nonpurified diet ad libitum will binge on a vegetable fat (shortening) when it is presented for 2 h each day, and this effect is enhanced when the fat is offered only 3 times per week. A similar finding has been reported with shortening that is trans-fat-free (18). Rats with restricted access to vegetable fat do not show alterations in body weight or body-fat accrual compared with nonpurified diet–fed controls (17,19); however, they do show elevated plasma leptin levels (19).

Models of bingeing on sweet-fat mixtures
The combination of sweet and fat activates multiple taste receptors, postingestive signals, and neuropeptide systems. Sugar-fat combinations, in the form of cookies or sugar-fat mixtures, have been used by Boggiano and others to induce binge eating in laboratory models (20,21). We have developed a model of binge eating using a nutritionally complete sweet-fat diet in rats that are not food-restricted (22). Rats with 2-h daily access to a sweet-fat food [Research Diets #12451 pellets, 45% fat, 20% protein, 35% carbohydrate, 4.7 kcal/g (20 kJ/g)] binge on it, even though they have access to standard rodent nonpurified diet ad libitum for the other 22 h/d. By wk 3 of access, the binging behavior is most pronounced, and these rats consume, on average, 58% of their daily energy intake during the 2-h period of access to the sweet-fat food (Fig. 2 A). These rats self-restrict their intake of standard nonpurified diet, similar to the effects we have reported with sugar (23) and others have reported with fat (17,19) or sugar-rich diets (14). Cyclic bingeing and self-imposed food restriction result in fluctuations in daily body weight characterized by weight loss between binges (Fig. 2 B). However, even if we take into consideration the self-restriction of standard rodent nonpurified diet between binges, an overall increase in body weight occurs in rats binging on sweet-fat pellets when compared with control groups that are fed only standard rodent nonpurified diet or access to the same sweet-fat pellets ad libitum (Fig. 2 C). Thus, this model represents binge eating that can result in increased body weight.

Food addiction
Many scientists have speculated that obesity and eating disorders, such as bulimia and anorexia, may have properties of an “addiction” (24–30). Moreover, several popular self-help books have been written on the topic of “sugar addiction” (31–34, to name just a few). Clinical and laboratory animal studies reveal similarities between overeating and drug addiction.

Clinical support for the theory of food addiction
A recent clinical study suggests that carbohydrates can have abuse potential for “carbohydrate cravers” (35). Likewise, craving-related changes in response to palatable foods have been identified using brain imaging techniques, and these changes are similar to those seen during drug craving (36,37). Dopamine (DA) has been suggested to have a common role in drug abuse.

Abbreviations used: ACh, acetylcholine; DA, dopamine; GAL, galanin; NAc, nucleus accumbens.
and obesity (28). Positron emission tomography scans reveal that obese subjects show a reduction in striatal D2 receptor availability that is correlated with the body weight of the subject (38) and similar in magnitude to the reductions reported in drug-addicted subjects (39). Opioids have also been the focus of clinical studies (25). Appetite dysfunctions in the form of either binge eating or self-starvation can affect endogenous opioid activity (40). Collectively, these clinical studies support the view that overeating can affect behavior and brain systems in a manner that resembles aspects of an addiction.

Behavioral evidence of sugar dependence in laboratory animals

Many of the behaviors and neurochemical changes that are characteristic of drug abuse are also apparent in our animal model of sugar bingeing described above and summarized in Table 1. This model is reviewed and related to the substance abuse literature in greater detail elsewhere (16).

Briefly, rats given daily intermittent access to a sugar solution and nonpurified diet escalate their sugar intake and increase their intake during the first hour of daily access, which we define as a “binge” (15). Sugar-bingeing rats show signs of opiate-like withdrawal when administered a relatively high dose of the opioid antagonist naloxone (3 mg/kg, subcutaneous). Somatic signs of withdrawal, such as teeth chattering, forepaw tremor, and head shakes, as well as behavioral manifestations of anxiety, are observed (41). Similar signs of opiate-like withdrawal emerge spontaneously without the use of an opioid antagonist when all food is removed for 24 h (23,41). Sugar-bingeing rats lever press for 23% more sugar in a test after 2 wk without sugar than they ever did before (42), suggesting a change in the motivational impact of

Table 1: Signs of dependence observed in sugar-bingeing rats

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<thead>
<tr>
<th>Sign</th>
<th>Result</th>
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<tr>
<td>Behavioral signs</td>
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<td>15, 55</td>
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<td>Sensitization</td>
<td>Large meals of sugar in the form of binges</td>
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<td>Opiate-like withdrawal</td>
<td>Anxiety, somatic indications</td>
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<td>Deprivation effect</td>
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<td>Locomotor sensitization to cocaine</td>
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<td>Dopamine</td>
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<td>Increased D1 receptor binding in the NAc</td>
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<td>Decreased D2 receptor binding in the</td>
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<td>Decreased D2 receptor binding in the NAc</td>
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<td></td>
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<td>Opioids</td>
<td>Respond to naloxone with signs</td>
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<td>Increased µ-opioid receptor binding in the</td>
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<td>Decreased enkephalin gene expression</td>
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1 Adapted with permission from Avena et al. (71). STR, striatum; DAT, dopamine transporter; VTA, ventral tegmental area.
sugar that persists and increases throughout a period of abstinence. We have also shown that rats bingeing on sugar develop locomotor cross-sensitization to a low challenge dose of amphetamine (0.5 mg/kg, intraperitoneally) that has little or no effect on naive rats (43). When rats are bingeing on sugar and then are forced to abstain, they subsequently show enhanced intake of 9% alcohol (44), suggesting that intermittent access to sugar can be a gateway to alcohol use.

Other researchers have obtained supportive behavioral findings using similar models of sugar bingeing. Signs of anxiety have been reported in rats with limited access to a high-sucrose diet (14). The mere removal of sugar has been reported to decrease body temperature (45). Also, aggressive behavior has been observed during removal of a diet that involves intermittent sugar access (46). Using operant conditioning, Grimm et al. (47) find that sucrose seeking increases during a month of sugar abstinence in rats that had intermittent sugar access. Intermittent sucrose access cross-sensitizes not only with amphetamine (43) but also with cocaine (48) and facilitates sensitization to the DA agonist quinpirole (49). These results support the theory that the DA system is sensitized by intermittent sugar access; this is important because enhanced mesolimbic dopaminergic neurotransmission plays a role in the behavioral effects of sensitization as well as cross-sensitization (50) and may contribute to addiction (51,52).

**Neurochemical evidence of sugar dependence**

The evidence described above suggests that sugar bingeing can produce behaviors that are similar to those observed in drug-dependent rats. Concomitant neurochemical changes may result in, or perpetuate, these behaviors. These signs are also summarized in Table 1 and are explained in greater detail in an earlier article (16).

We have found changes in DA, acetylcholine (ACh), and opioid systems in sugar-bingeing rats that are similar to those observed with some drugs of abuse. Autoradiography reveals increased D₁ receptor binding in the nucleus accumbens (NAc) and decreased D₂ receptor binding in the striatum relative to nonpurified diet-fed rats (15). Rats with intermittent sugar and nonpurified diet access also have decreased D₂ receptor mRNA in the NAc, and increased D₁ receptor mRNA in the NAc and dorsal striatum compared with nonpurified diet-fed controls (53). Sugar-bingeing rats have a significant decrease in enkephalin mRNA (53), whereas µ-opioid receptor binding is significantly enhanced in the NAc shell, cingulate, hippocampus, and locus coeruleus (15).

One of the strongest neurochemical commonalities between sugar binging and drugs of abuse is their effect on extracellular DA. A hallmark of drugs that are abused is repeated increase in extracellular DA, whereas during normal feeding, the DA response fades out after repeated exposure to a food (54). When rats are bingeing on sugar, the release of DA is recurrent, which may make the brain adapt as it does to a drug of abuse. Rats that are bingeing on sugar apparently release DA every day, as measured on d 1, 2, and 21 of access (55). Control rats fed sugar or nonpurified diet ad libitum, rats with intermittent access to just nonpurified diet, or rats that taste sugar only 2 times, develop a blunted DA response that is typical of a food that loses its novelty.

Withdrawal from drugs such as morphine, nicotine, and alcohol is often accompanied by alterations in DA/ACh balance in the NAc: specifically, DA decreases while ACh increases (56–58). Rats binging on sugar also show this neurochemical imbalance in DA/ACh during withdrawal. This result occurs both when rats are given naloxone to precipitate opiate-like withdrawal (41) or after 36 h of food deprivation (23).

Others have reported supportive findings. There is a decrease in D₂ receptor binding in the NAc of rats with intermittent access to sucrose and nonpurified diet compared with rats fed intermittent nonpurified diet only (59), and alterations occur in accumbens DA turnover and DA transporter binding in rats maintained on an intermittent sugar-feeding schedule (12,60).

**Is there evidence of dependence on fat or sweet-fat combinations?**

The literature suggests that, as with sugar, a similar addictive-like state may emerge with fat. Le Magnen (29) noted that naloxone could precipitate withdrawal in rats fed a cafeteria-style diet ad libitum that contains a variety of fat- and sugar-rich foods (e.g., cheese, cookies, chocolate chips). More recently, Teegarden and Bale (61) show that mice given access to diets high in fat or carbohydrate ad libitum for 4 wk and then forced to abstain endure an aversive environment to gain access to their preferred food. They conclude that withdrawal of such a diet elevates the stress state, contributing to dietary relapse. Also, Corwin and colleagues have shown an increase in progressive-ratio responding in rats that are binging on fat (62).

In terms of neurochemistry, it appears that binge eating of fat has effects on the accumbens DA and enkephalin systems that are similar to those observed with sugar bingeing. Limited exposure to fat (corn oil) will repeatedly release DA in the NAc, and this effect is caused by the taste of the oil (63). Rats with limited daily access to a sweet-fat diet show a significant decrease in enkephalin mRNA in the NAc (64), similar to the finding reported above with sugar (53). The role of opioids in the paraventricular nucleus of the hypothalamus has been studied using a binge model (65), and the findings suggest that d-Ala², NMe-Phe³, Gly-ol⁴-enkephalin increases fat intake in fat-prefering rats but has no effect in sucrose-prefering rats. These results indicate a complex role for paraventricular nuclear opioids in food intake, with preference and nutrient type affecting the ability of these compounds to change behavior.

Based on this neurochemistry and the behaviors described above, it seems logical that fat bingeing might also produce addictive-like behaviors. However, the data are not clear. Although fat offered ad libitum has been reported to produce some addictive-like behaviors (29,61), bingeing might enhance these effects. We have investigated whether behavioral signs of dependence emerge when animals binge using a variety of different high-fat diets and sweet-fat combinations. We have tested rats with limited (12-h or 2-h) access to a sweet-fat diet (Research Diets #12451, 45% fat, 20% protein, 35% carbohydrate), 12-h access to a sweet-fat mixture (35.7% vegetable fat, 64.3% sucrose), or 12-h access to vegetable fat (100% Crisco vegetable shortening), all with nonpurified diet concurrently available. Control groups were fed these diets ad libitum or given standard nonpurified diet ad libitum. After 21–25 d on the diets, rats were administered 3 mg/kg subcutaneous naloxone and then observed for somatic signs of distress and anxiety in the elevated plus maze. No significant evidence of opiate-like withdrawal was found with any of these fat-rich dietary options, in either the bingeing groups or those given food ad libitum, even though these procedures gave positive results in our previous reports with sugar bingeing (41). In other studies, we attempted to elicit signs of spontaneous opiate-like withdrawal by food depriving the rats maintained on fat-rich diets for 24–36 h. Again, although we report signs of anxiety and somatic indications of distress following fasting in sugar-bingeing rats (23), this was not observed in rats that had been bingeing with a high-fat source in the diet.
Although we have not noted signs of opiate-like withdrawal in fat-bingeing rats, that does not mean that excessive fat intake cannot produce addictive-like behaviors. Withdrawal is not a necessary criterion for drug craving, just as food deprivation is not necessary for food craving (37). Moreover, different classes of drugs (e.g., DA agonists, opiates) result in specific behavioral and physiological withdrawal signs. Thus, it may be that different macronutrients may also produce different withdrawal signs. It has yet to be determined whether or not binging on fat can precipitate other addictive-like behaviors, including cross-sensitization and abnormal motivation caused by abstinence.

Why do signs of opiate-like withdrawal emerge with sugar but not fat binging?

The relative lack of opiate-like withdrawal signs after fat binging underscores the importance of opioid systems in differentiating sugars and fats and their subsequent effects on behavior. The neuropeptide galanin (GAL) and its binding sites are expressed in brain areas important for both drug and food reward (11). GAL is considered a fat-stimulated peptide because its expression is increased in these brain regions in response to a high-fat meal (66). In addition, hypothalamic injection of GAL promotes the intake of fat in preference to carbohydrate in some situations (67,68). Interestingly, peripheral injection of galnon, a synthetic GAL agonist, decreases opiate withdrawal signs in morphine-dependent mice (69). A single systemic injection of galnon in GAL-knockout mice is sufficient to reverse some of the biochemical changes brought about by morphine administration (70). Thus, GAL may be an endogenous negative regulator of opiate reward by attenuating some of the behavioral and neurochemical effects of opiates. Based on these data, it is possible that that lack of opiate-like withdrawal signs in fat-bingeing rats may be caused by fat-induced endogenous GAL activation, which can inhibit the relevant opioid effects.

Implications for eating disorders and obesity

We began this article with a discussion relating binge eating to obesity. Indeed, the findings with animal models that have been presented suggest that binge eating of sugar, and possibly even fat, may have some addictive-like properties. However, sugar binging does not affect body weight, but a combination of sweet and fat does result in increased body weight (22). Thus, fat may be the macronutrient that results in excess body weight, and sweet taste may be largely responsible for producing addictive-like behaviors that include a withdrawal syndrome.

Other articles in this supplement include references (73–75).

Acknowledgment

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