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Running Head: FOOD CRAVING REAPPRAISAL

Neural systems underlying the reappraisal of personally-craved foods

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Abstract

Craving of unhealthy food is a common target of self-regulation, but the neural systems underlying this process are understudied. In the present study, participants used cognitive reappraisal to regulate their desire to consume idiosyncratically craved or not craved energydense foods, and neural activity during regulation was compared to each other and to activity during passive viewing of energy-dense foods. Regulation of both food types elicited activation in classic top-down self-regulation regions including the dorsolateral prefrontal, inferior frontal, and dorsal anterior cingulate cortices. This main effect of regulation was qualified by an interaction such that activation in these regions was significantly greater during reappraisal of craved (versus not craved) foods, and several regions, including the dorsolateral prefrontal, inferior frontal, medial frontal, and dorsal anterior cingulate cortices, were uniquely active during regulation of personally-craved foods. Body mass index significantly negatively correlated with regulation-related activation in the right dorsolateral prefrontal cortex, thalamus, and bilateral dorsal anterior cingulate cortex, and with activity in nucleus accumbens during passive viewing of craved (versus neutral, low energy density) foods. These results suggest that several of the brain regions involved in the self-regulation of food craving are similar to other kinds of affective self-regulation, and that others are sensitive to the self-relevance of the regulation target.

Overweight and obesity are at unprecedented levels. As such, gaining a better understanding of the psychological and neural determinants of eating behavior is a pressing topic for research. Food elicits strong appetitive and affective responses, which motivate us to consume it (Desmet & Schifferstein, 2008). In fact, the stimuli themselves don't even need to be present to elicit these responses; pictures of food elicit cravings and other affective responses (Heatherton & Wagner, 2011; Hofmann, et al., 2010). Advertisers capitalize on these responses, inundating potential customers with pictures of juicy hamburgers, crisp fries, and creamy chocolate desserts. While some of these advertised foods may be healthy, the billions of advertising dollars spent every year by the fast food industry are mainly focused on marketing unhealthy food (Harris, Schwartz, & Brownell, 2011). These dollars are not wasted; subjective cravings for energy-dense (ED) foods are associated with an attentional bias toward those foods (Papies, Stroebe, & Aarts, 2008; Smeets, Roefs, & Jansen, 2009), increased eating of ED foods (Gilhooly, et al., 2007), and future increases in weight (Yokum, Ng, & Stice, 2011).

People have the ability to modulate responses to these food stimuli using various forms of self-regulation. One kind of affective self-regulation is cognitive reappraisal, the reinterpretation of an emotional stimulus or event so as to change its meaning (Giuliani & Gross, 2009). Reappraisal is an effective way of modulating positive emotions in general (Giuliani, McRae, & Gross, 2008) and craving of preferred foods in particular (Giuliani, Calcott, & Berkman, 2013). Researchers have proposed that eating behavior is the result of an interplay between appetitive motivation and cognitive control processes such as reappraisal (Nederkoorn, et al., 2010). A new but substantial body of research has begun to uncover the brain systems involved in desire for and appetitive motivation toward food (e.g., Killgore, et al., 2003; Pelchat, et al., 2004; Siep, et al., 2009), primarily focusing on obesity (e.g., Carnell, et al., 2012; De Silva, et al., 2012).

However, the brain systems involved in regulating those responses in normal weight individuals have received far less attention.

Considerable knowledge about other forms of self-regulation can serve as a starting point for understanding food craving regulation. For example, deficits in self-regulation and executive function suffered by patients with frontotemporal dementia, which include binge eating and strong cravings for sweets and carbohydrates (Mendez, Licht, & Shapira, 2008), have been traced to atrophy of the frontal cortex (Piguet, 2011). While much of the research on the neuroscience of self-regulation in healthy adults has focused on the regulation of emotion and thought (Heatherton, 2011), work extending these findings to the regulation of a range of targets (e.g., food craving, affect, thought) suggest that, regardless of the regulation target, studies tend to reveal activation during self-regulation in the ventromedial and lateral prefrontal cortex (PFC), and the anterior cingulate cortex (ACC; Heatherton, 2011). A recent review of the brain regions engaged during emotion regulation using cognitive reappraisal found consistent recruitment of lateral PFC and ACC, but not ventromedial PFC (Buhle, et al., 2013). Of the two studies to date directly investigating the neural correlates of food craving reappraisal in healthy-weight individuals (Hollmann, et al., 2011; Siep, et al., 2012), only Hollmann et al. (2011) explicitly compared reappraisal with passive viewing. In that study, reappraisals focusing on the long-term negative consequences of consumption elicited strong activation in the PFC (dorsolateral prefrontal and inferior frontal cortices). Furthermore, no studies have examined whether regulation-related brain activity relates to real-world measures associated with food intake, such as body mass index, in healthy-weight individuals.

In light of the limited number of studies investigating the neural systems of food craving regulation, the present research aims to clarify how the brain supports the self-regulation of food craving using cognitive reappraisal, and whether and how activation in those neural systems is related to real-world measures related to food intake. Our study design has two unique features that advance the literature. First, we capitalize on the idiosyncratic nature of food preferences (Rozin & Vollmecke, 1986) to develop task conditions that enable comparisons that can be difficult to capture with other normed targets of regulation. Though some past studies of the reappraisal of negative emotion have created participant-specific stimuli (e.g., negative autobiographical memories; Goldin, et al., 2013), the conditions of the present paradigm allowed us to examine both the general neural systems of food craving regulation (with the main effect of self-regulation versus viewing), and also the neural systems that are specific to the selfregulation of idiosyncratically craved foods (with the simple effect of self-regulation of craved versus not craved foods). This second comparison is specifically designed to yield a more ecologically valid picture of how everyday regulation operates because people presumably dedicate more time and effort in their daily lives to regulating targets that are self-relevant (e.g., craved foods, personally emotional stimuli) compared to generic targets that are not necessarily self-relevant (e.g., ED foods that are not craved, emotional images from the International Affective Picture System). Second, we measured body mass index (BMI) as an index of cumulative energy-dense food intake. Weight gain in part results from an imbalance between the energy content of food eaten and the energy expended by the body (Bolton-Smith & Woodward, 1994; George, et al., 1990; Hall, et al., 2012), and thus BMI is expected to be related systematically (though not perfectly) to food intake. We argue that studying the relationship between BMI and neural activation is a logical first step toward understanding how the basic neurocognitive systems involved in cognitive self-regulation are related to behaviors that take place primarily outside the laboratory. As such, this study contributes to an emerging interest

within cognitive neuroscience about the links between neural functioning and real-world outcomes (Berkman & Falk, 2013).

We hypothesize that food craving regulation will elicit brain activation in the domaingeneral self- and emotion-regulation networks reviewed above, including the dorsolateral prefrontal cortex (DLPFC), inferior frontal gyrus (IFG), and dorsal ACC (dACC). Furthermore, we hypothesize that regulating responses to personally-craved ED foods (versus ED foods that are not personally-craved) will recruit additional regions beyond the typical self-regulation network—including the medial PFC, which is implicated in self-relevance processing (Kelley, et al., 2002)—but we note that there are no previous results directly relevant to this comparison on which to base a hypothesis. Demonstrating whether or not food craving reappraisal recruits these brain regions is important for several reasons. First, if we can verify that reappraisal of food craving relies upon similar neural regions as reappraisal of other kinds of emotional stimuli, then we could import knowledge gained from studies of emotion regulation into the field of craving regulation. Second, it will broaden the available options for studying reappraisal beyond the standard battery of negative pictures, which are not appropriate for many populations (e.g., children). We also expect that passive viewing of personally-craved ED foods (versus viewing of not craved ED foods) will elicit activation in the nucleus accumbens (NAcc), a hub of the dopaminergic reward system consistently identified in human and animal studies of reward, drug addiction, motivation, and food processing (Demos, Heatherton, & Kelley, 2012). Lastly, we hypothesize that activation in these regions will relate meaningfully to real-world measures associated with food self-regulation, including BMI and self-reported everyday food regulation. A mechanistic understanding of the self-regulation of appetitive responses to stimuli that are harmful in the long term has profound implications for both goal-directed food consumption

specifically and for cognitive and behavioral self-regulation broadly.

Methods

Participants

Fifty participants (17 male, age M = 21.77, SD = 2.36), recruited from the University of Oregon undergraduate and graduate student population, completed a study lasting approximately 90 minutes. Potential participants were excluded if they were left-handed, below 18 or above 30 years of age, non-native English speakers, had a current or past diagnosis of neurological or psychiatric disorder, had a history of head trauma, were pregnant, currently used psychoactive medication, had any non-MRI compatible conditions (e.g., metal in body), were currently on a diet, were on a diet in the past 3 months, or planned to go on a diet in the next 3 months. To ensure they were neither sated nor food deprived, participants were instructed to eat a full meal approximately 2 hours before the start of the scan. All gave informed consent in accordance with the University of Oregon Institutional Review Board. Two additional participants were excluded from analyses due to a history of brain trauma not disclosed during prescreening. Task

Details of the task are outlined in our previous work (Giuliani, et al., 2013). Briefly, images of two types of palatable foods were included as stimuli: low energy density foods ("Neutral"), and ED foods of the participants' choosing. The total stimulus set consisted of 20 pictures of low energy-density food (carrots, cauliflower, celery, corn, cucumber, beans, broccoli, Brussels sprouts, eggplant, lettuce, squash, tomatoes; pre-tested desirability M = 2.48, SD = .27), and 40 pictures in each of the following categories of ED food (pre-tested desirability Ms = 3.46-3.52, SDs = .17-.35): chocolate, cookies, donuts, fries, ice cream, pasta, and pizza. Importantly, images were chosen such that the mean desirability of the ED food categories were

not significantly different from each other (all paired-samples p-values > .05), and that the mean of each ED food category was significantly greater than the mean of the Neutral stimuli (pvalues < .001).

All participants saw the same set of 20 Neutral stimuli. For the ED stimuli, participants chose from the list of seven food types listed above the one that they craved the most ("Craved") and the one they craved the least ("Not Craved"), and saw only images within those two categories in addition to the Neutral stimuli during the task. Craving was defined as the desire and tendency to consume the target food, even in the absence of hunger. The ED food trials were organized in a 2 (Stimulus: Craved vs. Not Craved) x 2 (Instruction: Look vs. Regulate) design. The stimuli assigned to each condition were counterbalanced across participants, and the stimuli within each category did not vary from participant to participant. The final event-related design included 5 trial types (Look Neutral (LN), Look Craved (LC), Look Not Craved (LNC), Regulate Craved (RC), and Regulate Not Craved (RNC)), with 20 trials for each condition totaling 100 trials, distributed across three approximately even runs with brief breaks in between.

The Look instruction directed participants to focus on the pictured food, imagine it was actually in front of them, and think about consuming it. The Regulate instruction directed participants to focus on the food, imagine it was in front of them, and think about it in a way that reduced their desire to eat it. We asked participants to use one of the following four reappraisal strategies specific to food, developed on a separate sample (Giuliani et al., 2013): 1) imagine that you are currently very full, 2) focus on the negative consequences of eating the pictured food (e.g., stomachache, weight gain), 3) remind yourself that you can save the food for later, and 4) imagine that something bad had happened to the food (e.g., sneezed on). Participants chose one strategy before the task and were directed to use that same strategy on every trial. For the rating

period, we instructed participants to report their craving honestly at the end of each trial. To minimize the demand characteristics of the task regarding regulation success (i.e., reduced desire ratings on Regulate trials), we explicitly stated "we don't expect you to be able to do this on every picture, so please honestly rate how much you desire the food when all is said and done." To minimize instruction contamination across trials, we instructed participants to view each trial as a fresh event, and to do their best to only look or only regulate according to the cue.

Each trial began with a 2s instructional cue (Look or Regulate), followed by a 5s stimulus presentation, 4s to rate the desirability of the stimulus, and a jittered inter-trial-interval averaging 1s and following a gamma distribution. Desirability ratings ("How much do you desire to eat this food?") were made on a 1-to-5 Likert scale, where 1="not at all" and 5="very much." Within-run stimulus order was optimized to maximize contrast estimation efficiency using a genetic algorithm (Wager & Nichols, 2003). Run order was counterbalanced across subjects. Strategy training

Prior to beginning the task, participants underwent a structured training session in which they received the strategy instructions described above and viewed a sample trial for each of the two instructions. Sample trials provided participants experience with using the cognitive reappraisal strategies while looking directly at pictures of foods not used during the experimental session. The experiment began when the training session was complete and participants indicated to the experimenter that the directions and procedures were understood.

Experimental Procedure

After providing informed consent, participants were trained on the task, and then asked to choose their most and least craved categories of ED stimuli, and the regulation strategy they believed would be most effective for them during Regulate trials. Participants then practiced the

task with the experimenter (N.G.). In the MRI, participants first completed three runs of the eating regulation task, as well as other tasks not reported here. Following the completion of the task, the experimenter interviewed each participant to ensure that they had indeed used the selected regulation strategy on all Regulate trials. Lastly, participants completed the individual difference measures detailed below, reported height, weight, and level of hunger on a 1 ("very hungry") to 5 ("very full") Likert scale, and were thanked for their time.

Behavioral Data Analysis

Self-reported ratings of desire for the ED foods were subjected to a 2x2 repeatedmeasures analysis of variance to determine main effects of stimulus and instruction and test for interaction effects. Pairwise t-tests between conditions were performed to decompose observed effects, as well as investigate the differences between LN, LC, and LNC. Regulation success was defined in two ways: 1) percent difference in self-reported desire to consume the pictured food between Regulate and Look (i.e., the main effect of regulation) and 2) the percent difference in self-reported desire between LC to RC (i.e., the simple main effect of regulation of personallycraved foods). Reactivity to food cues was also defined in two ways: 1) percent difference in self-reported desire to consume the pictured food between LN and LC, and 2) percent difference in self-reported desire to consume the pictured food between LNC to LC. The alpha level was set to .05 for all analyses. All statistical analyses of behavioral data were performed in SPSS19.0 (IBM Corp., Armonk, NY).

Functional MRI Data Acquisition and Analysis

Data were acquired using a 3.0 Tesla Siemens Allegra head-only scanner at the University of Oregon's Robert and Beverly Lewis Center for Neuroimaging. Blood oxygen-level dependent echo-planar images (BOLD-EPI) were acquired with a T2*-weighted gradient echo

sequence (TR = 2000ms, TE = 30ms, flip angle = 80° , matrix size = 64×64 , 32 contiguous axial slices with interleaved acquisition, field of view = 200mm, slice thickness = 4mm). This sequence also prospectively corrected for motion during acquisition with PACE (Thesen et al., 2000). For each participant, a high-resolution structural T1-weighted 3D MP-RAGE pulse sequence (TR = 2500ms, TE = 4.38ms, TI = 1100ms, flip angle = 8° , matrix size = 256×192 , 160 contiguous axial slices, voxel size = 1mm³, slice thickness = 1mm; time 10:49) was acquired coplanar with the functional images. After the functional runs, field map scans were acquired to obtain magnetization values used to correct for field inhomogeneity (TR = 500ms, TE = 4.99ms, flip angle = 55°, matrix size = 64 x 64, field of view = 200mm, 32 contiguous axial slices with interleaved acquisition, slice thickness = 4mm).

Before preprocessing, non-brain tissue was removed from the brain images using robust skull stripping with the Brain Extraction Tool (BET) in FMRIB's Software Library (FSL; http://www.fmrib.ox.ac.uk/fsl/). Image preprocessing was implemented in SPM8 (Wellcome Department of Cognitive Neurology, London, UK; http://www.fil.ion.ucl.ac.uk/spm/), which included correction for field inhomogeneities, realignment and co-registration of functional images to each subject's own high-resolution structural image using a six-parameter rigid body transformation model, reorientation to the plane containing the anterior and posterior commissures, spatial normalization into space compatible with an MNI atlas, and smoothing using a 6mm³ full-width half-maximum Gaussian kernel. Statistical analyses were implemented in SPM8. For each subject, event-related condition effects were estimated according to the general linear model, using a canonical hemodynamic response function, high-pass filtering (128s) and a first-order autoregressive error structure. At the subject level, BOLD signal was modeled in a fixed effects analysis with separate regressors modeling each condition of interest

during the picture presentation period (5s) and for the instruction and rating periods. Linear contrasts were created for each comparison of interest (e.g., RC vs. LC). These contrasts were then imported to group-level random effects analyses for inference to the population.

Because the brain regions previously identified in reappraisal encompass several large cortical and subcortical regions, we investigated the neural correlates of food regulation using whole-brain analyses. For these whole-brain analyses, we applied a combined voxel-height and cluster-extent correction for multiple comparisons to guard against Type I error derived from AFNI's AlphaSim software (Cox, 1996). AlphaSim takes into account the size of the search space and the estimated smoothness of the data (using AFNI's 3dFWHMx) to generate probability estimates (using Monte-Carlo simulations) of a random field of noise producing a cluster of voxels of a given size for a set of voxels passing a given voxel-wise p-value threshold. In our data set, these simulations determined that a voxel-wise threshold of p < .001 combined with a spatial extent threshold of 25 voxels corresponded to a family-wise error (FWE) corrected false-positive probability of p < .05 across the whole brain. For large clusters identified in wholebrain analyses that encompassed multiple anatomical brain areas, we individually interrogated the peaks by creating regions of interest (ROI). Spherical (10mm) ROIs were built in MarsBar (MRC Cognition and Brain Sciences Unit, Cambridge, UK; http://marsbar.sourceforge.net/), centered at the peak voxel for each anatomically-defined region within the larger cluster.

Although the main focus of this study is food craving regulation, our design also allowed us to replicate the finding that the NAcc is involved in ED food reactivity. Due to the small volume of this region, we created a priori ROIs encompassing the left and right NAcc, as defined by the Harvard-Oxford subcortical atlas in FSL (left: 4280mm²; right: 4296mm²). These ROIs were then imported into MarsBar, where average ROI signal intensities (beta values) were

calculated separately for each condition compared with fixation baseline.

Individual Difference Measures

We obtained several measures to index the external validity of neural responses during the task. First, we calculated Body Mass Index (BMI; weight in kg/(height in m)²; M = 21.71, SD = 2.8, range = 17.48-31.63) using self-reported weight and height at time of scan. Because the distribution of BMI was significantly non-normal, a squared transformation was applied to improve normality. Second, although self-reported trait measures of regulation do not consistently correlate with task-based laboratory measures (McRae, 2013), we nonetheless administered questionnaires related to food craving regulation that may tap a different, more trait-like aspects of food-related self-regulation. Specifically, participants completed the Three-Factor Eating Questionnaire (TFEQ; Stunkard & Messick, 1985); cognitive restraint average M =.44, SD = .21, α = .82; disinhibition average M = .37, SD = .2, α = .75) and the Dutch Eating Behavior Questionnaire (DEBQ; van Strien, et al., 1986); restrained average M = 2.38, SD = .83; $\alpha = .92$; external average M = 3.33, SD = .59, $\alpha = .8$). Relationships between these measures and the beta values extracted from neural activation were calculated using Pearson's correlations in SPSS.

Results

Behavioral

Replicating our previous behavioral study with this task (Giuliani et al., 2013), we observed significant main effects of Instruction (Look M = 3.11, SD = .59; Regulate M = 2.06, SD = .49; $F_{(1.49)} = 106.31$, p < .001) and Stimulus (Craved M = 3.09, SD = .48; Not Craved M = 3.09; SD = .48; Not M = 3.09; SD = .48; Not M = 3.09; M = 32.08, SD = .55; $F_{(1.49)}$ = 115.05, p < .001) on self-reported desire to consume the depicted food. As shown in **Figure 1**, reappraisal successfully reduced self-reported desire to consume both the Craved $(M = 2.44, SD = .68; t_{(49)} = 11.02, p < .001)$ and Not Craved foods (M = 1.67, SD = .46; $t_{(49)} = 7.99$, p < .001) as compared to passive viewing (Look). This was qualified by a Stimulus by Instruction interaction ($F_{(1.49)} = 38.57$, p < .001), indicating that the magnitude of regulation success (reduction in self-reported desire) was greater for Craved than Not Craved foods.

Paired samples t-tests revealed that reactivity was greater than zero under both definitions (LN to LC, LNC to LC). LC foods (M = 3.75, SD = .6) were rated as significantly more desirable than LN $(M = 2.41, SD = .84; t_{(49)} = 10.03, p < .001)$ and LNC foods $(M = 2.48, SD = .81; t_{(49)} =$ 11.00, p < .001), and there was no significant difference in the rated desirability of the LN and LNC stimuli. Neither reactivity definition varied significantly by gender or hunger level.

Across all participants, 21 chose to imagine that something bad had happened to the pictured food (42%), 17 chose the "focus on the negative consequences of eating that food" strategy (34%), 11 chose the "imagine that you are currently very full" strategy (22%), and 1 chose the "remind yourself that you can save that food for later" strategy (2%). The distribution of participants selecting these strategies is not significantly different from that reported in our previous publication using this paradigm (Giuliani, et al., 2013). Regulation success (percent difference in self-reported desire between LC and RC) did not differ significantly by strategy used, gender, or self-reported general hunger of the participant.

fMRI

Regulation. As shown in Figure 2a, the main effect of Regulate versus Look across both picture types (Craved, Not Craved) produced a large cluster of activation encompassing the left IFG, anterior insula, and DLPFC, which connected with another cluster in the dACC and presupplementary motor area (pre-SMA). Other clusters included the left right inferior parietal lobe (IPL) and supramarginal gyrus, right anterior insula, bilateral thalamus, bilateral posterior

cingulate (PCC), and right DLPFC (see **Table 1**). The inverse contrast, the main effect of Look versus Regulate across both picture types, showed significant clusters of activation in the bilateral inferior occipital gyri, and the left posterior insula.

Further interrogation of the significant clusters from the Regulate > Look contrast broken down by condition produced two distinct activation patterns. First, the clusters located in the bilateral PCC, left and right MTG and thalamus, and left IPL and supramarginal gyrus showed the expected pattern of the Regulate > Look main effect in both the Craved and Not Craved conditions. Second, in the large cluster containing left IFG, anterior insula, DLPFC, dACC and pre-SMA, and the smaller clusters located in the right DLPFC and right anterior insula, activation during RC was significantly greater than any of the other four conditions, indicating a possible interaction between Stimulus and Instruction in these regions.

A whole-brain investigation of the voxels significantly more active in the RC > LC contrast as compared to the RNC > LNC contrast was conducted to formally test for the Stimulus by Instruction interaction. This contrast revealed additional clusters including the left and right medial superior frontal gyrus (SFG), right DLPFC, and right IPL (Figure 3, for full list see **Table 2**). To further understand the nature of this interaction, we individually interrogated the simple effects of RC > LC and RC > RNC. The whole-brain contrast of RC > LC (Figure 2b, **Table 2**) revealed significant clusters of activation in bilateral dACC and pre-SMA, left IFG into anterior insula and DLPFC, left IPL and supramarginal gyrus, right IFG and anterior insula, left MTG, right DLPFC, right STG, right MTG, right caudate, bilateral PCC, and right cerebellum. The whole-brain contrast of RC > RNC (**Table 2**) revealed significant clusters of activation in the left and right IPL, anterior insula and IFG, bilateral dACC extending into right SFG, bilateral PCC, left MFG, right DLPFC, and right SFG. None of the activation from these regulation

contrasts differed by reappraisal strategy, food category chosen, gender or hunger.

Reactivity. Both left and right NAcc were significantly more active during LC compared to LNC (left: t = 2.24, p = .015; right: t = 1.93, p = .029). Interestingly, activation in the left and right NAcc ROIs did not differ significantly between the LC and LN conditions, perhaps because of the heterogeneity of responses to the low energy density "Neutral" foods. We supplemented these a priori ROI analyses with exploratory whole-brain contrasts investigating both definitions of reactivity (**Table 3**; all at FWE p < .05). The contrast of LC > LNC revealed significant clusters in the IPL and lingual gyrus, left IPL and posterior insula, and the bilateral dACC and superior PCC¹. The contrast of LC > LN revealed three significant clusters of activation, in the right cuneus, precuneus, and parahippocampal gyrus. None of these contrasts varied significantly by food category chosen, gender, or hunger.

Relationships between neural activity and real world measures of regulation and reactivity

Regulation. BMI negatively correlated with RC vs. LC activity in the thalamus (r = -.31,p = .032), and the right DLFPC (r = -.29, p = .047; Figure 4). Within the peaks of the large PFC cluster, the right (x = 6, y = 20, z = 34; r = -.28, p = .049) and left dACC (x = -3, y = 23, z = 37; r = -.28= -.28, p = .053), BMI and regulation-related activity were negatively related such that BMI decreased as brain activity during regulation increased. The cognitive restraint subscale of the TFEQ and the restrained subscale of the DEBQ were used to assess the relationship between task-based neural and self-reported trait indices of food craving regulation. After multiple comparisons correction, regulation-related brain activity did not relate to either subscale.

Reactivity. Left and right NAcc ROI activity during reactivity operationalized as LC >

¹ Unsurprisingly, relaxing the cluster threshold reveals a small cluster in the left NAcc (t = 3.33; k = 2; peak at x = -12, y = 17, z = -8), an anatomical region small enough that we did not expect it to survive the 25 3mm³ voxel cluster threshold.

LNC did not relate significantly to the disinhibition subscale of the TFEQ, the external subscale of the DEBQ, self-reported reactivity (LC-LN, LC-LNC), or to BMI. However, similar to previous work (Demos et al., 2012), NAcc activity during reactivity operationalized as LC > LN positively correlated with BMI across all subjects (left: r = .33, p = .019; right: r = .43, p = .002).

A follow-up analysis of the relationships between reactivity- and regulation-related brain activity and BMI showed that regulation-related right DLPFC activation moderated the relationship between reactivity-related right NAcc activation and BMI ($t_{(1.46)} = 3.3$, p = .076), such that the positive relationship between BMI and right NAcc activity during reactivity was attenuated with increased right DLPFC activity during regulation.

Discussion

This study examined how self-regulation of craving for tempting foods influenced selfreported desire and neural activation. We capitalized on the idiosyncratic nature of food craving to characterize the brain systems specific to regulating personally-craved foods as well as those involved in regulating appetitive desires in general. Reappraisal effectively decreased selfreported desire for food and elicited activation in the left IFG and IPL, left and right DLPFC, anterior insula, and MTG, and bilateral thalamus, dACC, and PCC. Activation in a subset of those regions (DLPFC, IFG, and dACC) was specific to regulation of personally-craved foods. Finally, regulation-related activity in the thalamus, bilateral dACC, and right DLPFC was correlated with BMI, which is related to overall ED food intake (Bolton-Smith & Woodward, 1994; George, et al., 1990; Hall, et al., 2012).

For both self-reported reduction in food craving and reappraisal-related brain activation, there was a difference between regulation of idiosyncratically craved and not craved foods. Selfreport ratings revealed a significant stimulus type (Craved, Not Craved) by instruction (Look,

Regulate) interaction such that the regulation-related decrease in food desire was greater for the Craved versus Not Craved foods. Within the brain regions showing significantly more activity during regulation of ED foods versus passive viewing, some appeared to be regulation-general (e.g., similar levels of activation during both RC and RNC as compared to LC and LNC) whereas others showed significantly greater activation during RC trials compared to any other trial type. A direct comparison investigation of the stimulus by instruction interaction revealed that the left and right medial SFG, right DLPFC, IPL precentral gyrus, and precuneus displayed significantly more activation during the regulation of craving for Craved foods (RC > LC) compared to the regulation of craving for Not Craved foods (RNC > LNC). Within these regions, the left and right medial DLPFC, right SFG, right precentral gyrus were recruited by regulation of both targets but to a greater degree for the reappraisal of personally-craved foods, while the right IPL was uniquely recruited during craved food regulation. This supports our hypothesis that reappraising personally-craved foods recruits the medial PFC regions implicated in the processing of self-relevant information (Kelley, et al., 2002). Furthermore, it indicates that reappraisal of personally-relevant appetitive stimuli recruits an overlapping but distinct set of brain regions than general craving regulation, and suggests that regulation of idiosyncratic stimuli may be different than regulation of generic stimuli. The burgeoning field of food craving regulation has recognized the importance of using idiosyncratic stimuli (e.g., Hollmann et al., 2011), but studies of other types of self-regulation (e.g., emotion regulation) are just beginning to use this approach (e.g., Goldin et al., 2013). This emerging body of literature suggests that taking the self-relevance of the target into account is critical for future research on self-regulation. Indeed, it may be the case that the self-relevance of stimuli is a robust moderator of other effects studied in cognitive neuroscience more broadly. If so, the field could capitalize on self-relevance

by taking advantage of current advances in computer and information technology that allow scientists to build highly individualized simulations of real-world phenomena in the lab.

Interestingly, the brain regions classically recruited by cognitive reappraisal of negative emotional stimuli (Buhle, et al., 2013), including the dACC and DLPFC, were among those recruited to a greater degree by reappraisal of personally-craved foods. In other words, the data presented here clearly implicate these regions in the self-regulation of appetitive motives (e.g., craving). Taken together with previous data, this provides good evidence of the flexibility of those regions to regulate both approach and avoidance (e.g., fear, disgust) responses. While the activation in these regions was seen in the main effect, simple effect analyses revealed that they were driven by RC-related activation. We believe that these stimuli recruit these brain regions because of the high amount of conflict resolution (dACC) and top-down regulation (DLPFC, IFG) required to reappraise craved foods as compared to not craved foods. These findings have strong implications for food consumption interventions aimed at reducing the intake of energydense foods, as individuals who find this sort of direct top-down cognitive regulation of desire to be challenging may be directed to find other regulation strategies (e.g., avoiding the foods via situation selection or modification; Gross, 1998).

Reappraisal of both craved and not craved foods significantly reduced activation in the left and right occipital cortex, and the left posterior insula (Look > Regulate). While it is surprising that we did not find ventral striatum (e.g., NAcc) down-regulation during reappraisal, this null result is not unprecedented. For example, Hollmann et al. (2011) found no reduction in ventral striatum activity during regulation; in fact, they found increased dorsal striatum activity during the cognitive control of eating, which they interpreted to reflect increased integration of reward and goal-directed behavior. The relatively few extant fMRI studies on the cognitive

control of food craving make it challenging to know exactly what role the striatum plays in this process. One possibility is that null findings may be due to the relatively small anatomical size of the dorsal and ventral striatum. Another possibility, which warrants further study after two independent null findings, is that the field may need to reconsider the role of the ventral striatum in food craving regulation. It may be that ventral striatum responses to food cues are relatively immune to reappraisal, and that decreased reports of craving result solely from increased regulatory activity "overpowering" the reactivity response without decreasing it.

We observed increased activation in the occipital lobe and nucleus accumbens during the viewing of craved foods, which replicates previous work on food cue reactivity (Demos, et al., 2012; Schur, et al., 2009). Perhaps not surprisingly, the patterns of reactivity to craved foods differed depending on the comparison condition. Only the comparison of LC to LNC stimuli produced significantly greater activation in the NAcc, but this activation was not related to selfreported food cravings or weight. In contrast, comparing BOLD signal during viewing of craved foods to healthy (neutral) stimuli did not elicit significant activation in the NAcc, but the degree of activation NAcc in this contrast was significantly related to BMI. This relationship supports previous findings in the field, which documented the positive correlation between NAcc activation and food stimuli (Born, et al., 2011), as well as actual snack consumption (Lawrence, et al., 2012). These results indicate that both neutral and personally-undesired ED foods are useful control conditions because they may contain distinct information.

Given the high level of heterogeneity in eating behavior and food-related self-regulation across people, the dual-relationship between BMI with both regulation- and reactivity-related brain activation is of particular interest. First, although they were both correlated with BMI, regulation (DLPFC) and reactivity (NAcc) activity were not significantly related to each other.

This suggests that there may be at least two separable pathways through which food-related neural responses are related to cumulative food consumption measures such as BMI. Second, an exploratory analysis of these relationships showed that the positive relationship between BMI and right NAcc activity during reactivity was attenuated with increased right DLPFC activity during regulation. While the interpretability of this finding is limited by the trend-level significance of this finding and the self-reported nature of the height and weight values used to calculate BMI, this indicates that regulation-related activation of the DLPFC may be of particular interest to investigators developing targeted interventions, as it moderates the link between neural reactivity to craved food stimuli and BMI.

While brain activity during regulation and reactivity related meaningfully to BMI, we also did not see a relationship between task-based neural or behavioral indices of food regulation and self-reported trait food regulation as measured by the TFEQ and DEBQ. This uncoupling of regulation frequency and success has been documented in the literature (see McRae, 2013), and the present findings add to a growing body of research observing this effect. The present findings also challenge the restriction-based nature of current dieting interventions, which ask people to eat less energy-dense food but do not generally teach them how to mange their cravings for these popular foods. We suggest that affecting the antecedent mental processes associated with the consumption of energy-dense foods via reappraisal as may be more successful than simply targeting the consumption behavior itself. Furthermore, while BMI was used as a proxy for cumulative ED food intake, and related significantly to both food reactivity- and regulationrelated brain activity, more sensitive measures of body composition and eating behavior may produce more sensitive indices of these brain-behavior relationships. Third, while previous research has noted marked gender differences in eating behavior and body-weight regulation

(Cornier, et al., 2010; Rolls, Federoff, & Guthrie, 1991; Wang, et al., 2009), we did not find any significant gender effects in the present study. This may be because we selectively recruited participants who were not actively dieting, thereby removing known gender effects on food conflict and dieting (Rolls, et al., 1991). Lastly, personally-craved and not craved foods were identified at the category level, but not at the actual stimulus level. We accepted this limitation in order to equate the categories on normative ratings, but it is possible that the observed differences between craved and not craved foods would have been even more pronounced if participants were able to identify each image as idiosyncratically craved or not.

In the present study, we showed that reappraisal of energy-dense food stimuli recruits top-down control brain regions recruited by other forms of self-regulation. For many of these regions, including the DLPFC and dACC, this recruitment was driven by the reappraisal of idiosyncratically-craved foods, perhaps due to the greater cognitive control and conflict resolution required to successfully reduce the desire to consume these foods. The knowledge that these two regions are heavily recruited during this cognitive process may help us in the future to leverage interventions that we know target the functioning of these regions through training (see Schweizer, et al., 2013) and apply them to eating. Further, the present results have strong implications for stimulus choice in self-regulation paradigms, as they demonstrate that the selfrelevance of stimuli can significantly affect both the magnitude and pattern of associated neural activity. In other words, self-relevance may be a robust moderator not only of self-regulation of food craving but also of other effects studied in cognitive neuroscience. As such, self-relevance may be a double-edged sword: on one hand, as a nuisance variable, the degree to which an individual relates to a stimulus may artificially inflate correlations between neural responses to that stimulus and other variables like self-report; on the other hand, scientists may be able to

boost the power of small N studies by leveraging the increased effect sizes that might be associated with tailored stimuli. The present findings answer calls for a greater breadth of experimental paradigms to study emotion regulation generally (Carter, 2009) and also test the range of the flexibility of the neural circuits involved therein to target different motivational and affective responses. Both reactivity-related activation in the NAcc and reappraisal-related activation in the DLPFC, thalamus, and dACC significantly related to BMI, suggesting that real world patterns of food consumption, which influence body weight, may result from both reactivity to and regulation of food cues. Indeed, the present findings indicate that a neurallyinformed intervention that targets both systems may indeed be the most effective means of reducing the consumption of energy-dense food.

Figure 1. Self-reported mean desire to consume pictured food in the five conditions: look neutral, look craved, look not craved, regulate craved, regulate not craved. Error bars represent SEM.

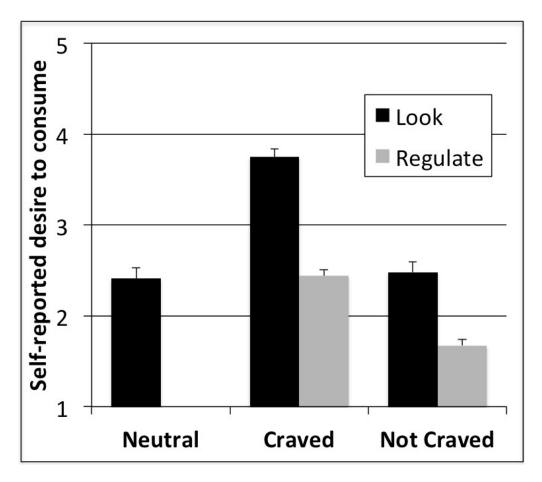


Figure 2. a) Main effect of Regulate (Regulate Craved + Regulate Not Craved) > Look (Look Craved + Look Not Craved), p < .001 (k = 25), and b) a visualization of the overlap of this contrast with Regulate Craved > Look Craved (RC > LC; p < .001, k = 25). Voxels only active in the Regulate > Look contrast are in yellow, only RC > LC are in red, and overlapping voxels are in orange.

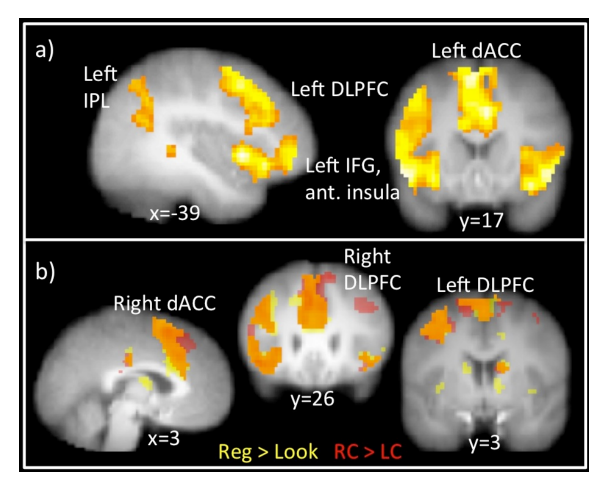


Figure 3. Stimulus x Instruction interaction (p < .001, k = 25) with bar graphs demonstrating the pattern of activation by condition for the left medial superior frontal gyrus (SFG), right dorsolateral prefrontal cortex (DLPFC), and right inferior parietal lobule (IPL).

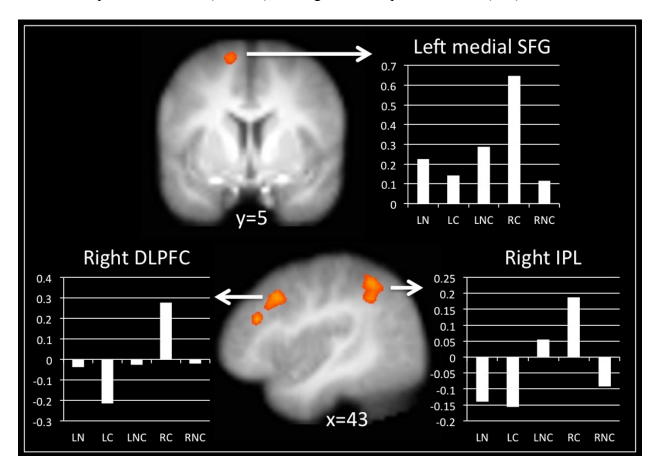


Figure 4. Scatterplot demonstrating the inverse relationship between Regulate Craved > Look Craved (RC > LC) activation in the right dorsolateral prefrontal cortex (DLPFC; peak at 42, 17, 43; square root transformation to improve normality) and body mass index (BMI; square transformation and z-scored).

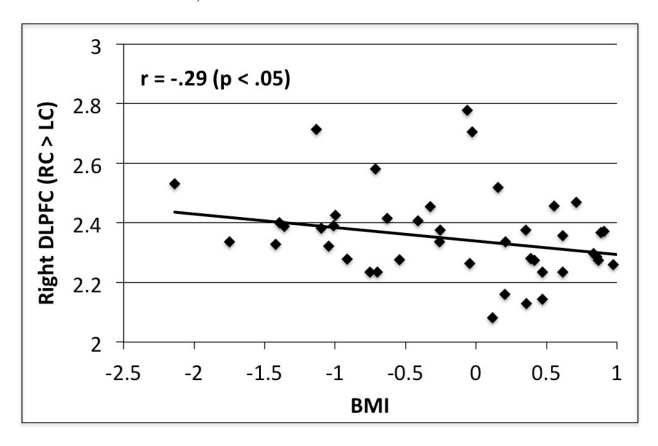


Table 1. Regions, MNI coordinates, cluster sizes and peak T-values for the Regulate > Look and Look > Regulate main effects (FWE p < .05 threshold used for all contrasts).

Contrast and Region	MNI Coordinates	Cluster size	Peak T		
	(x, y, z)				
Regulate (Craved + Not Craved) > Look (Craved + Not Craved)					
Left IFG	(-51, 11, -2)	2917	8.02		
Left anterior insula	(-30, 20, -11)		7.2		
Left DLPFC	(-36, 5, 58)		7.1		
Left supramarginal gyrus	(-51, -49, 31)	415	7.58		
Left IPL	(-51, -55, 43)		6.87		
Right anterior insula	(42, 11, -5)	435	7.21		
Right cerebellum	(33, -61, -35)	76	7.09		
Left middle temporal gyrus	(-57, -31, -8)	127	5.98		
Bilateral thalamus	(0, -7, 13)	210	5.69		
Right middle temporal gyrus	(48, -31, -11)	105	5.49		
Bilateral PCC	(-3, -22, 25)	65	5.01		
Right DLPFC	(36, 47, 19)	72	4.74		
Look (Craved + Not Craved) > Regulate (Craved + Not Craved)					
Right inferior occipital	(24, -88, -2)	721	6.84		
Left inferior occipital	(-18, -94, -8)	284	5.84		
Left posterior insula	(-48, -16, 16)	44	4.24		

Note: $DLPFC = dorsolateral\ prefrontal\ cortex;\ IFG = inferior\ frontal\ gyrus;\ IPL = inferior$ parietal lobe; PCC = posterior cingulate cortex.

Table 2. Regions, MNI coordinates, cluster sizes and peak T-values for the Stimulus x Instruction, Regulate Craved > Look Craved, and Regulate Craved > Regulate Not Craved contrasts (FWE p < .05 threshold used for all contrasts).

Contrast and Region	MNI Coordinates (x, y, z)	Cluster size	Peak T
Stimulus x Instruction			
Right medial SFG ^{^+}	(6, 29, 58)	99	5.33
Right DLPFC^+	(39, 20, 34)	154	5.22
Right IPL ⁺	(39, -52, 52)	171	5.04
Right precentral gyrus^+	(33, -4, 64)	30	4.46
Right precuneus	(36, -67, 43)	38	4.41
Left medial SFG ^{^+}	(-12, 5, 64)	38	4.4
Regulate Craved > Look Craved		•	
Bilateral dACC	(-3, 14, 52)	1065	8.24
Bilateral SFG	(0, 17, 61)		8.21
Left IFG	(-45, 23, -8)	1435	7.82
Left DLPFC	(-45, 2, 49)		7.61
Left IPL	(-51, -55, 43)	405	7.41
Left supramarginal gyrus	(-57, -52, 31)		6.72
Right anterior insula	(36, 11, -11)	463	7.2
Right IFG	(45, 20, -11)		6.2
Right cerebellum	(33, -58, -35)	78	6.79
Left middle temporal gyrus	(-60, -34, -5)	106	5.99
Right DLPFC	(42, 17, 43)	240	5.17
Right superior temporal gyrus	(54, -52, 34)	69	4.71
Right middle temporal gyrus	(45, -37, -5)	92	4.57
Right caudate	(15, -4, 16)	37	4.42
Bilateral PCC	(0, -22, 31)	42	4.36
Regulate Craved > Regulate Not Cr	aved		
Right IPL	(39, -43, 49)	278	5.77
Bilateral dACC into right SFG	(6, 29, 58)	573	5.41
Right anterior insula	(42, 20, 1)	289	4.96
Right IFG	(57, 14, 7)		4.9
Right DLPFC	(42, 20, 43)	126	4.91
Left MFG (pre-SMA)	(-39, -4, 49)	94	4.81
Left anterior insula	(-36, 17, 4)	54	4.64
Left IFG	(-42, 20, -11)		3.76
Right SFG	(12, 50, 40)	28	4.52
Bilateral PCC	(3, -22, 31)	45	4.39
Left IPL	(-33, -49, 46)	47	4.16

Note: dACC = dorsal anterior cingulate cortex; DLPFC = dorsolateral prefrontal cortex; IFG = inferior frontal gyrus; IPL = inferior parietal lobe; MFG = middle frontal gyrus; PCC =

posterior cingulate cortex; SFG = superior frontal gyrus; SMA = supplementary motor area. ^ denotes regions from the Stimulus x Instruction interaction that also showed a significant difference in the RC > LC simple contrast.

+ denotes regions from the Stimulus x Instruction interaction that also showed a significant difference in the RC > RNC simple contrast.

Table 3. Regions, MNI coordinates, cluster sizes and peak T-values for the Look Craved > Look Neutral and Look Craved > Look Not Craved contrasts (FWE p < .05 threshold used for all contrasts).

Contrast and Region	Coordinates	Cluster size	Peak T
	(x, y, z)		
Look Craved > Look Neutral			
Right cuneus	(15, -94, 7)	46	4.82
Right precuneus	(27, -64, 43)	92	4.64
Right middle occipital gyrus	(36, -76, 16)	25	3.97
Right parahippocampal gyrus	(27, -46, -20)	37	3.88
Look Craved > Look Not Craved			
Right IPL	(54, -37, 31)	47	4.99
Right lingual gyrus	(12, -70, -5)	42	4.54
Bilateral dACC	(3, 11, 31)	48	4.43
Left IPL/posterior insula	(-48, -22, 19)	92	4.37
Bilateral superior PCC	(3, -4, 46)	39	4.06

Note: dACC = dorsal anterior cingulate cortex; IPL = inferior parietal lobe; PCC = posterior cingulate cortex.

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