Effects of distraction on taste-related neural processing: a cross-sectional fMRI study

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ABSTRACT

Background: In the current obesogenic environment we often eat while electronic devices, such as smart phones, computers, or the television, distract us. Such “distracted eating” is associated with increased food intake and overweight. However, the underlying neurocognitive mechanisms of this phenomenon are unknown.

Objective: Our aim was to elucidate these mechanisms by investigating whether distraction attenuates processing in the primary and secondary taste cortices, located in the insula and orbitofrontal cortex (OFC), respectively.

Methods: Forty-one healthy, normal-weight participants received fixed amounts of higher- and lower-sweetness isocaloric chocolate milk while performing a high- or low-distracting detection task during fMRI in 2 test sessions. Subsequently, we measured ad libitum food intake.

Results: As expected, a primary taste cortex region in the right insula responded more to the sweeter drink ($P < 0.001$, uncorrected). Distraction did not affect this insular sweetness response across the group, but did weaken sweetness-related connectivity of this region to a secondary taste region in the right OFC ($P$–family-wise error, small-volume corrected $= 0.020$). Moreover, individual differences in distraction-related attenuation of taste activation in the insula predicted increased subsequent ad libitum food intake after distraction ($r = 0.36$).

Conclusions: These results reveal a mechanism explaining how distraction during consumption attenuates neural taste processing. Moreover, our study shows that such distraction-induced decreases in neural taste processing contribute to individual differences in the susceptibility for overeating. Thus, being mindful about the taste of food during consumption could perhaps be part of successful prevention and treatment of overweight and obesity, which should be further tested in these target groups. This study was preregistered at the Open Science Framework as https://bit.ly/31RtDHZ. 

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Introduction

Worldwide, the prevalence of obesity has nearly tripled since 1975. In 2016, >1.9 billion adults were overweight, with 650 million being clinically obese (1). The problem of obesity has been partly attributed to the obesogenic food environment, which offers an enormous variety of palatable, energy-dense, easily consumed foods (2, 3). Furthermore, people’s lifestyles have changed over the last decades, with increasing demands of multitasking due to their interaction with electronic devices [e.g., televisions, computers, and smart phones (4)]. As a consequence, people often eat while engaged in activities that prevent them from focusing on satiation signals such as sensory stimulation from the food products they are consuming or gastric signals [e.g., (2, 5)]. Such “mindless” or distracted eating has been causally related to increased immediate and later food intake and is associated with increases in BMI (2, 6–10).

However, the underlying neurocognitive mechanism of how distracted eating could increase food intake remains elusive. In rodents, others have found that distraction decreases and slows down processing of taste information in the taste cortex (11). In humans, it has been suggested that distraction attenuates taste
FIGURE 1  Timeline of the experimental sessions. After a screening session, participants came to the laboratory twice, for 2 experimental sessions. Except for the difference in attentional load (high or low) during the distraction task, the 2 sessions were identical. Session order was counterbalanced across participants. Between t₀ and t₃₀, participants performed the high- or low-distraction task during fMRI scanning. In each session, participants performed 80 trials (4 blocks of 20 trials). To manipulate distraction, 90% of trials were of low load (high-frequency trials) and 10% of high load (low-frequency trials) in the low-load session, and vice versa for the high-load session. Each block had 8 trials of low sweetness, 8 of high sweetness, and 4 of neutral taste. After the task, participants were removed from the MR scanner and watched a documentary in the behavioral laboratory until t₇₅. Subsequently, participants consumed a chocolate snack ad libitum. Glucose (glu) measurements and VAS hunger, fullness, thirst, ideal sweetness, liking, sweet and savory desire, nausea, and anxiety were rated at several time points. See the Methods section for further details. MR, magnetic resonance; VAS, visual analog scale.
FIGURE 2 Trial structure of the categorical visual detection task. Each trial started with an instruction screen, indicating the target category (furniture, tool, or toys) and attentional load (low "->" or high ">>>") of the trial. Then, pictures were presented followed by a visual mask, and subjects were instructed to push a button as fast as possible upon detection of pictures belonging to the instructed category. During each trial, participants were administered a fixed amount of lower- or higher-sweetened chocolate milk, or a tasteless neutral solution through a gustometer. Markers were placed on the participant’s neck to enable detection of participants’ swallow movements. Onsets and offsets of the swallow movements were used to determine trial durations in the first level (single-subject) fMRI models. ITI, inter-trial interval.

abstain from eating solid foods and from drinking sugared or sweetened drinks (but not water) 6 h prior to the experiment and to refrain from alcohol use (24 h) and neuroleptic or psychotropic drug use (7 d). To standardize participants’ hunger, they were instructed to ingest a standardized load 3 h before each session (yogurt drink, strawberry flavor: 200 g; 850 kJ, 6.0 g protein, 30.0 g carbohydrates, 6.0 g fat; Breaker; Melkunie).

At the start of the first test session, anthropometric measurements were taken. Subsequently, participants underwent an fMRI scan for 30 min in which they performed a categorical visual detection task (Figure 1 and Figure 2). During the task, participants received higher-sweetness (120 g) or lower-sweetness (120 g) chocolate milk, or a tasteless solution (60 g), through small tubes. After scanning, participants watched a documentary (BBC Life, “Primates or Plants”; order was randomized). Subsequently, participants were seated in front of a bowl with colored button-shaped chocolates (M&Ms; Mars Wrigley) for 10 min, and were asked to eat until comfortably full. In session 1, the participants completed the Behavioural Inhibition System/Behaviour Approach System (22), the Baratt Impulsiveness Scale-11 (23), and Kirby (delayed reward discounting) (24) questionnaires. In session 2, they completed the following questionnaires: the Binge Eating Scale (25), Food Frequency Questionnaire–Dutch Healthy Diet (26), Dutch Eating Behaviour Questionnaire (27), and the Power of Food Scale (28), which are used to describe the study population (see Supplemental Table 1 for mean scores and SDs).

During the test session, glucose measurements were taken through finger pricks and analyzed with use of a glucose meter (Stat Strip Xpress®; Nova Biomedical). This was done at 4 time points (t): t0min (baseline, before first chocolate milk exposure), t50min (directly following last exposure to chocolate milk), t50min+ and t75min (before consumption of the chocolate snack).

At several time points during the experiment, participants rated how hungry, full, and thirsty they were on a paper (t-10min and t75min) or digital (t0min, t30min, t50min, and t75min) version of a visual analog scale (VAS; 100-mm line), with anchors ranging from 0 (“not at all”) to 10 (“very”). Before and after the task (t0min and t50min), participants also digitally rated how nauseous and anxious they felt, and their desire for something savory and something sweet.

At least 1 wk after their first test session (mean difference ± SD: 11.93 ± 7.92 d), participants revisited the laboratory for their second test session. The time of day at which participants had their experimental sessions varied between subjects (minimum: 10:45 h; maximum: 19:10 h). Within-subject, we aimed to plan a participant’s second session at a similar time of day as his/her first session (mean ± SD time difference: 1.00 ± 1.52 h).

Gustatory stimulation
Gustatory stimuli were determined in a pilot study (see Supplemental Methods: Gustatory pilot study). The higher- and lower-sweetness chocolate milk were solutions of cocoa powder (Blooker; Bickery Food Group; 2 g), dextrine-maltose (Fantomalt; Nutricia; 9 g) in whole milk (3.5% fat/100 g), and
liquid noncaloric sweetener (Natrena; Jacobs Douwe Egberts B.V.; lower sweetness: 0.0867 g; higher sweetness: 1.5457 g/100 g whole milk). The neutral solution was determined based on work by Veldhuizen et al. (29), and contained 1.25 mol/m³ sodium bicarbonate and 12.5 mol/m³ potassium chloride in water.

Before and after the task that participants performed in the magnetic resonance (MR) scanner, they received 2 rounds of 3.75 g of the neutral drink and of the lower- and higher-sweetness drink for tasting through tubes innervated by pumps (gustometer; Watson-Marlow). For all drinks, they rated how much they liked the drink on VASs. For the 2 chocolate milk drinks, they also rated to what extent the sweetness of the chocolate milk matched their ideal sweetness on an ideal point scale.

Categorical visual detection task

Participants performed a categorical visual detection task during 3T fMRI scanning (Figure 2). The task was programmed with Presentation software (version 16; Neurobiobehavioral Systems, Inc.). Each trial (total duration: 12 s) started with a fixation cross (duration, 2 s), followed by an instruction screen (1 s), indicating the category of pictures to which the participant needed to respond (furniture, tools, or toys) and the speed of the trial (“fo r s l o wt r i a l ,” for a slow trial, “>>>” for a fast trial). For example, if the instruction screen stated “category: furniture, >>>>>,” this meant that participants needed to respond to stimuli in the category of furniture, and the pictures would be presented at high speed. In order to keep visual stimulation equal for both trial types, a visual mask always followed a picture. The visual masks were scrambled versions of the stimulus pictures, to keep luminance equal. For the low-speed trials, both pictures and visual masks were presented for 750 ms. For the high-speed trials, pictures were presented for 75 ms and the visual mask for 675 ms. Consequently, there were twice as many pictures and visual masks in the high-speed trials relative to the low-speed trials (12 vs. 6), thus a higher attentional load. Whenever a target stimulus was presented (i.e., a picture belonging to the instructed category), participants had to push a button upon detection with their right index finger. Participants made responses using an MRI-compatible button box. Participants received no feedback on whether they responded correctly.

During the trials, participants received a fixed amount (3.75 g) of chocolate milk of higher or lower sweetness or a fixed amount of the neutral solution through a gustometer. Drink administration started together with presentation of the first picture and lasted for 6 s (i.e., a flow rate of 0.625 g/s). The drinks were administered at room temperature. A dot changing in color in the center of the screen informed participants of the start (brown color) and finish (white color; 1 s) of administration. At the end of each trial, the dot turned green (2 s), cueing participants to swallow the milk or tasteless solution. As swallowing can also be an uncontrollable, reflexive movement, participants’ swallowing was filmed. A marker was placed on the Adam’s apple, as this area shows the most swallowed-related movement. Frame-by-frame video analysis of the marker’s movement was later performed to pinpoint the exact moments in time when participants swallowed during the experiment (see Supplemental Methods: Video analysis of swallow movements).

Participants performed 4 blocks of 20 trials (a total of 80 trials; see Figure 1). For the low-distraction session, 90% of the trials were low-speed trials (high-frequency trials: pictures presented at a slow pace), and 10% of the trials were high-speed trials (low-frequency trials: pictures presented at a fast pace). Thus, in the low-distraction session, each block contained 18 low-speed trials and 2 high-speed trials. For the high-distraction session, this division was the same, although in the opposite direction [90% high-difficulty trials (high-frequency trials), 10% low-difficulty trials (low-frequency trials)]. The low-frequency trials were added to ensure participants would keep anticipating the occurrence of both high- and low-speed trials in both sessions. Each block of 20 trials had 4 neutral taste trials. Trials 1, 7, 14, and 20 were always neutral. Of the remaining trials, 7 trials (50%) were of higher sweetness and 7 (50%) of lower sweetness. Category (whether participants had to respond to pictures in the category tools, toys, or furniture) and drink sweetness presentation were pseudo-randomized (i.e., the same category and sweetness were never presented >3 times in a row). Moreover, maximally 2 target stimuli (i.e., pictures belonging to the instructed category) were presented after another.

MRI data acquisition and analysis

To measure BOLD contrast, whole-brain functional images were acquired on a Siemens 3T Skyra MRI scanner (Siemens Medical Systems) using a 32-channel coil. During the task, 3D echo planar imaging scans using a T2*-weighted gradient echo multi-echo sequence (30) were acquired [voxel size: 3.5 × 3.5 × 3 mm isotropic, TR (repetition time) = 2070 ms, TE (echo time) = 9 ms; 19.25 ms; 29.5 ms; 39.75 ms; field of view (FoV) = 224 mm]. The slab positioning and rotation (average angle of 14° to the anterior commissure axis) optimally covered both prefrontal and deep brain regions. Before the acquisition of functional images, a high-resolution anatomical scan was acquired [T1-weighted magnetization-prepared rapid gradient-echo (MPRAGE) sequence: voxel size 1 × 1 × 1 mm, TR 2300 ms, TE 3.03 ms, 192 sagittal slices, flip angle 8°, FoV 256 mm].

fMRI data were preprocessed using SPM8 (www.filion.ucl.ac.uk/spm) and FSL version 5.0.11 (http://www.fmrib.ox.ac.uk/fsl/). The volumes for each echo time were realigned to correct for motion artefacts (estimation of the realignment parameters is done for the first echo and then copied to the other echoes). The 4 echo images were combined into a single MR volume based on 30 volumes acquired before the actual experiment started using an optimized echo-weighting method (30). Combined functional images were slice-time corrected by realigning the time series for each voxel temporally to acquisition of the middle slice and spatially smoothed using an isotropic 8-mm full-width at half-maximum Gaussian kernel. Next, Independent Component Analysis based Automated Removal of Motion Artifacts (ICA-AROMA) (31) was used to reduce motion-induced signal variations in the fMRI data. Subject-specific structural and functional data were then coregistered to a standard structural or functional stereotactic space [Montreal Neurological Institute (MNI) template], respectively. After segmentation of the structural images using a unified segmentation approach, structural images were spatially coregistered to the mean of the functional images. The resulting transformation matrix of the segmentation was then used to normalize the anatomical and

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Functional images were resampled at voxel size $2 \times 2 \times 2$ mm.

**Statistical analysis**

To test for pre-experimental differences in liking of the higher- compared with the lower-sweetness chocolate milk, we performed a repeated-measures ANOVA (IBM SPSS Statistics 23) with within-subject factor drink sweetness (lower sweetness, higher sweetness) on the mean baseline ratings.

We used the sensitivity index $d'$ to calculate participants’ task performance (see Supplemental Methods “$d'$ prime” for how this score was calculated). Mean $d'$ scores on the detection task were analyzed using repeated-measures ANOVA with attentional load (low, high) and drink type (lower-, higher-sweetness milk, neutral drink) as within-subject factors. Low-frequency conditions (i.e., 10% low-speed trials on the high-distraction session and vice versa) were excluded from this analysis, as the low number of trials in these conditions would likely bias the results.

Statistical analysis of fMRI data was performed using a general linear model approach in SPM8 (www.fil.ion.ucl.ac.uk/spm). The images of both experimental runs were combined into 1 model including the low- and high-distraction test sessions. At the individual (first) level, subject-specific data were analyzed using a fixed-effects model, which included 10 regressors of interest. For each experimental run, the first 4 regressors reflected the trials of low attentional load, lower-sweetness drink; low attentional load, higher-sweetness drink; high attentional load, lower-sweetness drink; and high attentional load, higher-sweetness drink. The fifth regressor reflected trials in which the neutral solution was given, which was always of high-frequency load, so of low attentional load in the low-distraction session and of high attentional load in the high-distraction session to rinse in-between chocolate milk trials. Durations reflected the moment the gustometer started drink administration until the first swallow of the participants, or in case these video data were not available, the moment the swallow was cued in the task (dot turning green, mean ± SD duration: 11.22 ± 0.80 s). All regressors were convolved with the canonical hemodynamic response function. Parametric modulators reflecting the number of button presses per trial were added to the model for each regressor of interest, to correct for signal change induced by the difference in number of targets between the low and high attentional load condition. High-pass filtering (128 s) was applied to the time series of the functional images to remove low-frequency drifts, and correction for serial correlations was done using a first-order autoregressive (AR-1) model. Signal variation in white matter and cerebrospinal fluid regions was also included.

At the group (second) level, we assessed the effect of load (high > low attentional load) in 2 ways. First, we contrasted high- with low-load trials by including the high-frequency trials of each test session only, meaning a contrast (comparison) between sessions (high-distraction session: high-load regressors across drink types > low-distraction session: low-load regressors across drink types). Second, we also added the low-frequency regressors to assess this contrast (high- + low-distraction session: high-load regressors across drink types > low-load regressors across drink types).

Taste-related brain areas (sensitive to sweetness) were localized with the contrast higher > lower sweetness ($P < 0.001$, uncorrected), over both test sessions. For this contrast, within-test session data were available; therefore, we did not make a second contrast including low-frequency regressors. We used the Automated Anatomical Labeling (AAL) atlas (32) to determine whether the areas activated in this functional contrast overlapped with the anatomical insula (bilateral insula) and OFC (bilateral superior, medial, mid-, and inferior orbitofrontal regions of this atlas).

To investigate the effect of attentional load on processing in primary and secondary taste-related areas, we assessed the interaction effect of load (higher > lower load) × sweetness (higher > lower sweetness) using the test sessions in which the load conditions were the high-frequency load condition. The activated taste-related regions (determined by the contrast higher > lower sweetness) were used as regions-of-interest (ROIs) for the load × sweetness interaction contrast. Mean parameter estimates (i.e., $\beta$ weights) were extracted from all voxels in both ROIs separately using MarsBar (33). The regionally averaged parameter estimates were analyzed using ANOVA with the same factors as in the whole-brain analyses. As 2 ROIs were tested, effects were considered significant when reaching a threshold of $P < 0.025$ (Bonferroni-corrected for multiple comparisons). In exploratory analysis, we also assessed the interaction effect of load and sweetness at the whole-brain level (on a $P < 0.001$ uncorrected and a whole-brain $P$ family-wise error (FWE) cluster < 0.05 corrected threshold).

As a secondary analysis, we performed a generalized psychophysiological interaction (gPPI) (34) analysis to investigate distraction-related differences in functional connectivity for taste-related processing. As a seed, we used a taste-related region from our localizer approach described above. To estimate the neural activity producing the physiological effect in the seed region for each subject, the BOLD signal was extracted from this region and deconvolved (35). This was included in the model as the physiological regressor. The durations for each of the relevant task conditions—1) low-load, lower-sweetness drink; 2) low-load, higher-sweetness drink; 3) high-load, lower-sweetness drink; and 4) high-load, higher-sweetness drink—were included as psychological regressors. The psychophysiological interaction was entered by multiplying the estimated neural activity in the seed region by the duration times for each of the task conditions, separately convolved with the hemodynamic response function, resulting in 9 regressors of interest on the first level (i.e., 1 physiological, 4 psychological, and 4 interaction regressors). For each subject, we created a psychophysiological interaction (PPI) contrast for the interaction effect of distraction (high > low attentional load) and sweetness (higher > lower sweetness). On the second level, this PPI contrast was analyzed separately using a 1-sample $t$ test. Statistical inference ($P_{\text{FWE}} < 0.05$) was performed at the cluster level, correcting for multiple comparisons over the a priori–defined small-search volume: bilateral insula and OFC [AAL atlas (32)].

The results of all random-effects fMRI analyses had a threshold of $P < 0.001$ (uncorrected) and statistical inference was performed at the cluster level, FWE-corrected ($P_{\text{FWE}} < 0.05$) for multiple comparisons over the search volume (the whole-brain or a priori–defined small-search volumes).

The effect of attentional load on subsequent ad libitum food intake was tested with a paired-samples $t$ test to compare the total...
amount of chocolate snacks consumed (grams) between the high- and low-distraction test sessions.

Blood-glucose concentrations were analyzed using the linear mixed model [lmer4, version 1.1-14 (36); package in R, version 3.5.1; https://www.r-project.org], because we expected variation across participants in the glucose response over time, and traditional repeated-measures ANOVA cannot account for such variation. Participants’ glucose concentrations were analyzed with time ($t = 0$ min, $t = 30$ min, $t = 50$ min, and $t = 75$ min) and load (low, high) as fixed factors. We used random intercepts for participants, as we expected fasted glucose concentrations to vary between participants. Moreover, we used random slopes for the time predictor to account for the expected variation across participants in glucose response over time.

To test whether mean appetite (hunger, fullness) and thirst ratings varied as a function of load, we used repeated-measures ANOVA with within-subject factors load and time (digital: $t_0$, $t_5$, $t_{10}$, $t_{30}$, paper: $t_5$, $t_{75}$).

In an exploratory analysis, we investigated whether ad libitum food intake, blood glucose concentrations, and appetite and thirst measures (filled out digitally or on paper) covaried significantly with the effect of load on processing in the taste-related areas differentially activated in the higher > lower sweetness contrast. For ad libitum food intake, a repeated-measures ANCOVA (IBM SPSS Statistics 23) was executed with load (high, low) and sweetness (higher, lower) as within-subject factors and the difference in food intake between the low- and high-distraction sessions as a covariate. For the blood glucose measurements, we performed an ANCOVA with glucose as a dependent variable and load (high, low) and time ($t_0$, $t_{30}$, $t_{50}$, $t_{75}$) as within-subject factors and the difference in food intake between the low- and high-distraction sessions as a covariate. For the blood glucose measurements, we performed an ANCOVA with glucose as a dependent variable and load (high, low) and time ($t_0$, $t_{30}$, $t_{50}$, $t_{75}$) as within-subject factors and the difference in food intake between the low- and high-distraction sessions as a covariate. For the blood glucose measurements, we performed an ANCOVA with glucose as a dependent variable and load (high, low) and time ($t_0$, $t_{30}$, $t_{50}$, $t_{75}$) as within-subject factors and the difference in food intake between the low- and high-distraction sessions as a covariate. For the blood glucose measurements, we performed an ANCOVA with glucose as a dependent variable and load (high, low) and time ($t_0$, $t_{30}$, $t_{50}$, $t_{75}$) as within-subject factors and the difference in food intake between the low- and high-distraction sessions as a covariate.

Next, we localized brain regions responding to the difference in sweetness of the chocolate milk (higher > lower sweetness; $P < 0.001$, uncorrected). As expected, within our a priori–defined search volume (i.e., the anatomically defined bilateral insula + bilateral OFC from the AAL atlas), clusters in the left and right insula responded to this difference (Table 1 and Figure 3). As the left cluster comprised only 2 voxels, further analyses focused on the right functional insula cluster for the effects of distraction. We did not find differential sweetness responses in the OFC.

Finally, we tested the effect of distraction (attentional load) on activation of the right insula cluster that responded to the sweetness manipulation. We found no evidence for this effect, as the averaged extracted condition parameter estimates across the right insula cluster did not show a load × sweetness interaction [$F(1,40) < 1$, $P = 0.711$]. We obtained similar results when we explored the interaction effect of load and sweetness at the whole-brain level, where we did not find any whole-brain–corrected results ($P_{FWE} < 0.05$; see Supplemental Figure 2 for the results at $P < 0.001$, uncorrected). However, when we used this right insula cluster that showed greater responses for high > low sweetness (at $P < 0.001$, uncorrected) as a seed in the functional connectivity (i.e., gPPI) analysis, it showed decreased connectivity with a region in the right OFC under high compared with low distraction during processing of sweet taste (higher > lower sweetness; Table 1 and Figure 4). This right OFC region (MNI-coordinates x, y, z (mm): 32, 28, −18; AAL: right inferior frontal gyrus, pars orbitalis) was significant within our a priori–defined
search volume of the insula plus OFC \( P_{\text{FWE}} \) (cluster, after small volume correction) = 0.020, \( t = 5.27, k = 116 \). This shows that distraction weakens functional connectivity between the right insula and right OFC during taste processing.

**Effect of distraction on appetite and thirst ratings**

Hunger, fullness, and thirst ratings that were filled out digitally during the task \((t_6, t_{5}, t_{10}, t_{30})\) and on paper before and after the test session \((t_5, t_{75})\) were analyzed separately (Table 2). All ratings showed main effects of time, except for hunger, which was not significant in the paper ratings. These effects indicate significant increases in fullness and significant decreases in thirst and hunger over the time course of the task (digital) or test session (paper), as anticipated. No effects of distraction (i.e., attentional load) on any of the ratings were found. In addition, we found no significant pre-experimental differences between the low- and high-distraction session for all self-reported measures.

**TABLE 1** Summary of brain regions exhibiting effects of distraction (attentional load), sweetness, interactions between distraction and sweetness, and the result of the connectivity analysis (gPPI)

| Label | Part of cortex (F, P, T, O, I, C) | MNI coordinates \( x, y, z, \text{mm} \) | Size, no. of voxels | \( P_{\text{FWE}} \) (cluster-level) | \( r \) value (peak) |
|-------|----------------------------------|---------------------------------|-------------------|-------------------------------|------------------|
| Effect of distraction (high > low attentional load; low-frequency regressors included)\(^2\) |  |  |  |  |  |
| Calcarine | L | O | \(-8\) | \(-100\) | \(2\) | 2532 | <0.001 | 8.57 |
| Calcarine | R | O | \(14\) | \(-92\) | \(0\) |  |  | 8.22 |
| Cerebellum | R | C | \(8\) | \(-78\) | \(-16\) |  |  | 8.06 |
| Superior motor area | L | F | \(-4\) | \(10\) | \(52\) | 2183 | <0.001 | 7.89 |
| Frontal medial lobe | L | F | \(-30\) | \(-4\) | \(52\) |  |  | 7.13 |
| Mid cingulum | R | F | \(6\) | 0 | \(46\) |  |  | 6.72 |
| Parietal superior lobe | R | P | \(24\) | \(-56\) | \(50\) | 376 | 0.003 | 7.63 |
| Precentral | R | F | \(42\) | \(4\) | \(30\) | 1177 | <0.001 | 7.28 |
| Frontal medial lobe | R | P | \(30\) | \(4\) | \(60\) |  |  | 6.21 |
| Precentral | R | F | \(42\) | \(0\) | \(46\) |  |  | 5.65 |
| Temporal medial lobe | R | T | \(44\) | \(-66\) | \(8\) | 788 | <0.001 | 7.21 |
| Medial occipital lobe | R | O | \(32\) | \(-70\) | \(22\) |  |  | 5.37 |
| Cerebellum | L | C | \(-38\) | \(-56\) | \(-34\) | 444 | 0.001 | 7.14 |
| Insula | L | I | \(-32\) | \(22\) | \(4\) | 1106 | <0.001 | 6.91 |
| Superior frontal gyrus, orbital | R | F | \(48\) | \(26\) | \(-6\) |  |  | 6.33 |
| Inferior frontal gyrus, triangular | R | F | \(58\) | \(22\) | \(8\) |  |  | 3.93 |
| Medial occipital lobe | L | O | \(-42\) | \(-72\) | \(6\) | 625 | <0.001 | 6.82 |
| Parietal superior lobe | L | P | \(-20\) | \(-62\) | \(52\) | 228 | 0.028 | 5.93 |
| Insula | L | I | \(-30\) | \(24\) | \(0\) | 466 | 0.001 | 5.66 |
| Insula | L | I | \(-34\) | \(20\) | \(10\) |  |  | 4.80 |
| Insula | L | I | \(-32\) | \(30\) | \(10\) |  |  | 4.74 |
| Effect of distraction (high > low attentional load; low-frequency regressors excluded)\(^2\) |  |  |  |  |  |
| Temporal medial lobe | R | T | \(54\) | \(-62\) | \(10\) | 210 | 0.043 | 4.37 |
| Calcarine | L | O | \(2\) | \(-88\) | \(-6\) | 350 | 0.005 | 4.37 |
| Effect of distraction (low > high attentional load; low-frequency regressors included)\(^2\) |  |  |  |  |  |
| Inferior occipital lobe | R | O | \(38\) | \(-88\) | \(-10\) | 6611 | <0.001 | 8.70 |
| Cuneus | R | O | \(10\) | \(-70\) | \(22\) |  |  | 8.66 |
| Calcarine | R | O | \(20\) | \(-64\) | \(10\) |  |  | 8.18 |
| Postcentral | R | P | \(60\) | \(-8\) | \(26\) | 3074 | <0.001 | 7.78 |
| Rolandic operculum | R | F, P | \(48\) | \(-6\) | \(20\) |  |  | 7.73 |
| Postcentral | R | P | \(56\) | \(-12\) | \(34\) |  |  | 7.58 |
| Fusiform area | L | T, O | \(-36\) | \(-82\) | \(-12\) | 693 | <0.001 | 7.36 |
| Medial occipital lobe | L | O | \(-36\) | \(-92\) | \(-6\) |  |  | 6.28 |
| Postcentral | L | P | \(-60\) | \(-16\) | \(38\) | 3082 | <0.001 | 7.31 |
| Parietal inferior lobe | L | P | \(-52\) | \(-34\) | \(44\) |  |  | 7.21 |
| Postcentral | L | P | \(-54\) | \(-14\) | \(32\) |  |  | 7.04 |
| Sweetness localizer, higher > lower sweetness, masked with AAL insula + OFC\(^3\) |  |  |  |  |  |
| Insula | L | I | \(38\) | \(-4\) | \(10\) | 12 | 0.373 | 4.69 |
| Insula | L | I | \(-34\) | \(-6\) | \(14\) | 2 | 0.565 | 3.37 |
| gPPI—interaction effect [distraction (low > high) > sweetness (higher > lower)]\(^4\) |  |  |  |  |  |
| Inferior frontal gyrus, orbital | R | F | \(32\) | \(28\) | \(-18\) | 116 | 0.020 | 5.27 |

\(^1\) For the main effects of distraction, results are shown for the comparisons including and excluding the low-frequency regressors (see Methods). AAL, Automated Anatomical Labeling; C, cerebellar cortex; F, frontal cortex; FWE, family-wise error; gPPI, generalized psychophysiological interaction; I, insular cortex; L, left; MNI, Montreal Neurological Institute; O, occipital (visual) cortex; OFC, orbitofrontal cortex; P, parietal cortex; R, right; T, temporal cortex.

\(^2\) \( P < 0.05 \), whole-brain FWE corrected.

\(^3\) \( P < 0.001 \), uncorrected. Areas showing overlap with a priori-defined AAL atlas regions.

\(^4\) \( P < 0.05 \), small volume, FWE corrected.
Distraction and taste processing

FIGURE 3  BOLD responses to the effect of sweetness. The left (2 voxels) and right (12 voxels) middle insula show larger responses to the higher- vs. lower-sweetness drink comparison (circled in blue; n = 41). Statistical parametric maps had a threshold of P < 0.001 uncorrected for visualization purposes. All statistical parametric maps were overlaid onto a T1-weighted canonical image. Slice coordinates are defined in MNI152 space and images are shown in neurological convention (left = left). MNI, Montreal Neurological Institute.

Blood glucose concentrations

For 4 participants, one of the glucose measurements failed. Therefore, these measurements were excluded from the analyses. Analysis of the blood glucose concentrations revealed a main effect of time \[t(1, 88.85) = 11.28, P < 0.001; \text{Supplemental Figure 3}\], meaning that participants’ blood glucose concentrations increased significantly over the 4 time points during and after chocolate milk consumption, as expected \[\text{mean (SE): } t_{0} = 4.43 (0.46) \text{ mmol/L}, t_{30} = 4.84 (0.74) \text{ mmol/L}, t_{50} = 7.14 (1.23) \text{ mmol/L}, t_{75} = 7.17 (1.29) \text{ mmol/L}\]. Furthermore, distraction (attentional load) did not affect these glucose increases over time \[t(1, 260.19) = 1.81, P = 0.072\]. However, an exploratory analysis did reveal a significant distraction-related decreased rise in glucose concentrations at \( t = 75 \) relative to baseline \[t(1, 260.19) = 1.81, P = 0.072\]. However, an exploratory analysis did reveal a significant distraction-related decreased rise in glucose concentrations at \( t = 75 \) relative to baseline \[t(1, 260.19) = 1.81, P = 0.072\]. Nevertheless, further post hoc analyses (see Supplemental Results: Blood glucose concentrations) showed that these small effects were not statistically reliable. Given the low physiological relevance, these findings are not further discussed.

Food intake

Chocolate snack intake after the scan session did not differ between the test sessions \[\text{mean (SEM): low load } = 65.6 (5.9) \text{ g, high load } = 68.1 (6.8) \text{ g}; t(1, 40) = -0.61; P = 0.546\]. However, further analyses showed a significant interaction between attentional load and session order (low-distraction session first, high-distraction session first) for snack intake \[F(1, 39) = 8.27, P = 0.007\]. Specifically, food intake was significantly higher in the second, relative to the first, test session, independent of the attentional load of the test session \[\text{mean (SEM): session 1 } = 61.2 (6.2) \text{ g, session 2 } = 72.4 (6.4) \text{ g}; F(1, 40) = 8.68; P = 0.005; \text{Supplemental Figure 4A}; see Supplemental Figure 4B for a replication of this effect in a pilot study described in the Supplemental Methods: Behavioural pilot study\]. It is likely that the interaction between load and session order for food intake is driven by a repetition effect. Given session order effects on food intake, session order was added as a between-subject factor to all other analyses (i.e., fMRI, glucose, and behavioral). None of the other reported results changed after correction for order, or interacted with order (all \( P > 0.1 \)).

FIGURE 4  Results of the gPPI analysis with the right insula seed region in the top left (in blue, extracted from the higher- vs. lower-sweetness comparison). Shown is the right orbitofrontal area exhibiting significantly \((P = 0.020 \text{ after SVC})\) higher sweetness-related functional connectivity with the seed region under low, relative to high, distraction. n = 41; gPPI, generalized psychophysiological interaction.
Discussion

Distracted eating has been convincingly associated with increased food intake (6), but the underlying neurocognitive mechanisms remained elusive. Here, we examined how distraction—operationalized by varying attentional load—affects taste activation in, and connectivity between, primary and secondary taste-related brain areas (insula and OFC, respectively), blood glucose concentrations, subsequent chocolate snack intake, and appetite and thirst ratings.

As expected, higher compared with lower sweetness elicited differential BOLD responses in the insula: a small (2-voxel) cluster in the left insula and a larger cluster (12 voxels) in the right middle insula. We found no differential responses to sweetness in the OFC. These findings fit with recent work by Dalenberg et al. (14), who showed that the insular cortex processes the presence, pleasantness, and concentration of taste. More specifically, the right insular cortex dominated processing of taste concentration (intensity) signals, whereas the left insular cortex was more involved in representation of the presence of a taste stimulus and its pleasantness. This is in line with results from 2 other studies that also showed lateralization of intensity to the right insula (15, 16). Some studies have related activity of the right insula to processing of pleasantness; however, they did not control for effects of intensity (21, 39). In the present study, we varied taste intensity by comparing activation in response to a higher-sweetness versus a lower-sweetness drink and corrected for subjective liking differences of the 2 drinks at baseline. Crucially, this correction did not change the results of the higher > lower sweetness contrast. Therefore, the currently observed responses of the insula in the higher > lower sweetness comparison were mainly driven by differences in intensity, explaining the dominance of the right insula.

Importantly, we demonstrated that distraction attenuated taste-related functional connectivity between the right insula—found for the higher > lower sweetness contrast—and an area in the OFC. This OFC region is located in the caudomedial OFC (18), which is thought to be a relay between the anterior insula and the caudolateral OFC that is responsive to the pleasantness of taste (18). Another study manipulated pleasantness of chocolate milk and tomato juice through satiation and showed that pleasantness of the drinks correlated with taste activation in the left and right OFC, with the latter overlapping with the region found in our study (40). Thus, our findings suggest diminished functional coupling between primary and secondary taste cortices by distraction.

Our results further indicate that some individuals were more sensitive to distraction-related attenuation of taste-related activation in the right insula than others. Only when high attentional load affected taste processing of the lower-sweetness drink in the insula did subsequent food intake of participants increase. Previous work also showed large variation between participants in the effects of distraction on food intake (41, 42). Furthermore, the meta-analysis by Robinson et al. (6) showed that specifically highly disinhibited eaters are less likely to decrease their food intake after or during distraction. The interindividual variability found in our study is therefore not surprising and future studies should further investigate what drives these individual differences.

One study also investigated effects of cognitive load on food-related processing during fMRI [(13); note, however, that these authors manipulated working memory instead of attentional load]. In that study, higher cognitive load diminished nucleus accumbens responses during categorization of high- versus low-calorie food pictures. In addition, they showed that cognitive load altered the functional coupling between the nucleus accumbens

![Table 2: Appetite (hunger, fullness) and thirst ratings, filled out digitally and on paper, averaged over distraction (high, low attentional load)](https://doi.org/10.1093/ajcn/nqaa118)
Mean parameter estimates are presented in arbitrary units (a.u.), ad libitum food intake in amount consumed in grams.

lower, but not the higher, sweetness drink in the high- but not low-distraction session seems to predict increased food intake in the high-distraction session. Correlations for the low (LDS; \( n \)) and ad libitum intake (low–high distraction session); \( r = 0.36, P = 0.023 \). (B) Separate correlations for the low (LDS; \( r = -0.01, P = 0.973 \)) and high (HDS) distraction session (\( r = 0.45, P = 0.004; \) higher–lower sweetness). (C) Correlations in the high-distraction session only, for lower (LS; \( r = -0.30, P = 0.056 \)) and higher (HS; \( r = -0.04, P = 0.823 \)) sweetness separately. Less activation for the lower, but not the higher, sweetness drink in the high- but low-distraction session seems to predict increased food intake in the high-distraction session. Mean parameter estimates are presented in arbitrary units (a.u.), ad libitum food intake in amount consumed in grams.

and the right dorsolateral prefrontal cortex for high- versus low-calorie food pictures (13). However, these authors studied cognitive load effects on hedonic brain responses in the nucleus accumbens during categorization of high- and low-calorie food pictures versus object pictures as edible or inedible, in the absence of consumption during the task. By assessing actual taste processing during consumption of drinks in the scanner, we add to these previous findings that distraction attenuates connectivity in the taste network, and that the attenuating effect of attentional load on taste-related processing in the insula predicts subsequent food intake.

We found that responses for the lower-sweetness drink, in particular, predicted increases in later food intake under high load. Interestingly, a study by Hoffmann-Hensel et al. (43) found a similar effect when assessing the impact of cognitive load on fMRI responses to low- and high-calorie food odors with the same working memory load manipulation as by van der Wal and van Dillen (12). Their behavioral data revealed diminished perceived intensity for low- but not high-calorie food odors during high cognitive load. Similarly, higher cognitive load decreased OFC responses for low-calorie, but not the high-calorie, food odors. Their results could be explained by higher saliency of the high-calorie food odors and this may apply to our findings as well; as the higher-sweetness drink is more salient compared with the lower-sweetness drink due to its higher sweetness, attentional load might less easily suppress higher sweetness taste perception. It is, however, important to note that olfaction and gustation follow similar, but not identical, neural pathways [e.g., (43, 44)]. Nevertheless, the current and previous findings suggest that the saliency of odors and tastes plays a role in how distraction affects processing of food-related stimuli.

One limitation of our study is that the design was optimized for the primary outcome measure (i.e., the fMRI effects). Therefore, the distraction manipulation had to be relatively subtle (i.e., only varying the speed at which pictures were presented). Distractions such as watching a movie versus doing nothing in the MR scanner provide a less-controlled fMRI comparison with additional sources of variation (e.g., in perceptual, mood, and memory processes) than just varying attentional load. The relative subtleness of the distraction manipulation could explain why we did not find group effects of attentional load on food intake, or on appetite and thirst ratings.

In conclusion, by using fMRI during consumption, we found that distraction reduced functional connectivity between taste-processing areas and that distraction-related attenuation of taste-related processing in the insula predicted subsequent food intake. This provides a neurocognitive mechanism that improves our understanding of (the susceptibility for) overeating, and points to a potentially important role for undisrupted taste processing in overeating. A better understanding is essential for the successful prevention and treatment of overweight and obesity, where being mindful about the taste of food during consumption could perhaps be part of the solution. In our current—overly distracting—society, attentive eating might be more important than ever, to protect taste processing from being disrupted. Future studies should investigate the role of attention in neural taste processing in obesity.

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References

1. World Health Organization. Obesity and overweight [Internet]. [cited 2019 Oct 2]. Available from: https://www.who.int/news-room/fact-sheets/detail/obesity-and-overweight.
2. de Graaf C, Kok FJ. Slow food, fast food and the control of food intake. Nat Rev Endocrinol 2010;6:290–3.
3. Rolls BJ. Plenary Lecture 1: dietary strategies for the prevention and treatment of obesity. Proc Nutr Soc 2010;69:70–9.
4. Currier LM, Rosen LD, Cheever NA, Lim AF. Causes, effects, and practicalities of everyday multitasking. Dev Rev 2015;35:64–78.
5. Gore SA, Foster JA, DiLillo VG, Kirk K, Smith WD. Television viewing and snacking. Eat Behav 2003;4:399–405.
6. Robinson E, Aveyard P, Daley A, Jolly K, Lewis A, Lyczett D, Higgs S. Eating attentively: a systematic review and meta-analysis of the effect of food intake memory and awareness on eating. Am J Clin Nutr [Internet] 2013;97:728–42. [cited in 2019]. Available from: http://ajcn.nutrition.org/content/97/4/728.full.pdf+html?mCht p=p10vidsp.ovidweb.cgi?T =J&JAPAGE =reference&D =e med11&NEWS =N&AN =2013291485.
7. Bolhuis DP, Lakemond CMM, de Wijk RA, Luning PA, de Graaf C. Multiple lifestyle and overweight among children aged 9–11 years: results from the UK site of the International Study of Childhood Obesity, Lifestyle, and the Environment. BMJ Open 2016;6(2):e010677.
8. Wiggers JL, Scherjon SA, van den Bergh LM, van de Heyning PH, van Strien T, et al. The Development and Field Validation of the Dutch Eating Behavior Questionnaire. Int J Eat Disord 2009;42:473–81.
9. Bickham DS, Blood EA, Walls CE, Shrier LA, Rich M. Characteristics of screen media use associated with higher BMI in young adolescents. Pediatrics [Internet] 2013;131:935–41. [cited in 2019]. Available from: http://pediatrics.aappublications.org/cgi/doi/10.1542/peds.2012 1197.
10. Willke HJ, Standage M, Gillison FB, Cumming SP, Katmarzyk PT. Television viewing and overweight and obesity among children aged 9–11 years: results from the UK site of the International Study of Childhood Obesity, Lifestyle, and the Environment. BMJ Open 2016;6(2):e010677.
11. Mazzucato L, La Camera G, Fontanini A. Expectation-induced modulation of metastable activity underlies faster coding of sensory stimuli. Nat Neurosci [Internet] 2019;22:787–96. [cited in 2019]. Available from: https://doi.org/10.1038/s41593-019-0364-9.
12. van der Wal RC, van Dillen LF. Leaving a flat taste in your mouth: task load reduces taste perception. Psychol Sci 2013;24:1277–84.
13. van Dillen LF, van Steenbergen H. Tuning down the hedonic brain: cognitive load reduces neural responses to high-calorie food pictures in the nucleus accumbens. Cogn Affect Behav Neurosci 2018;18:447–59.
14. Dahlenberg JR, Hoogeveen HR, Renken RJ, Langers DRM, ter Horst GJ. Functional specialization of the male insula during taste perception. Neuroimage [Internet] 2015;119:210–20. [cited in 2019]. Available from: http://dx.doi.org/10.1016/j.neuroimage.2015.06.062.
15. Smith DM, Gregory MD, Mak YE, Gitelman D, Mesulam MM, Parrish T. Dissociation of neural representation of intensity and affective valuation in human gustation. Neuroimage 2003;39:701–11.
16. Spetter MS, Smeets PAM, de Graaf C, Viergever MA. Representation of sweet and salty taste intensity in the brain. Chem Senses 2010;35:831–40.
17. Rolls ET, Yaxley S, Sienkiewicz ZJ. Gustatory responses of single neurons in the caudolateral orbitofrontal cortex of the macaque monkey. J Neurophysiol [Internet] 1990;64:1055–66. [cited in 1999]. Available from: http://www.physiology.org/doi/10.1152/jn.1990.64.4.1055.
18. Small DM, Bender G, Veldhuizen MG, Rudenga K, Nachtigal D, Felsld J. The role of the human orbitofrontal cortex in taste and flavor processing. Ann N Y Acad Sci 2007;1121:136–51.
19. Haase L, Cerf-Ducastel B, Murphy C. Cortical activation in response to pure taste stimuli during the physiological states of hunger and satiety. Neuroimage 2009;44:1008–21.
20. Small DM. Taste representation in the human insula. Brain Struct Funct 2010;214(5–6):551–61.
21. Small DM, Zatorre RJ, Dagher A, Evans AC, Jones-Gotman M. Changes in brain activity related to eating chocolate. Brain 2001;124:1720–26.
22. Curver CS, White TL. Behavioral inhibition, behavioral activation, and affective responses to impending reward and punishment: The BIS/BAS Scales. J Pers Soc Psychol 1994;67:319–33.
distractibility, and memory for advertisements. Am J Clin Nutr 2009;89:37–44.

42. Bellisle F, Dalix AM, Airinei G, Hercberg S, Péneau S. Influence of dietary restraint and environmental factors on meal size in normal-weight women: a laboratory study. Appetite 2009;53:309–13.

43. Hoffmann-Hensel SM, Sijben R, Rodriguez-Raecke R, Freiher J. Cognitive load alters neuronal processing of food odors. Chem Senses 2017;42:723–36.

44. Small DM, Zald DH, Jones-Gotman M, Zatorre RJ, Pardo JV, Frey S, Petrides M. Human cortical gustatory areas: a review of functional neuroimaging data. Neuroreport 1999;10:7–14.