

Effect of flavor on neuronal responses of the hypothalamus and ventral tegmental area

Van Opstal, A.M.; Van den Berg-Huysmans, A.A.; Hoeksma, M.; Blonk, C.; Pijl, H.; Rombouts, S.A.R.B.; Van der Grond, J.

Citation

Van Opstal, A. M., Van den Berg-Huysmans, A. A., Hoeksma, M., Blonk, C., Pijl, H., Rombouts, S. A. R. B., & Van der Grond, J. (2019). Effect of flavor on neuronal responses of the hypothalamus and ventral tegmental area. *Scientific Reports*, *9*, e11250. doi:10.1038/s41598-019-47771-8

Version: Publisher's Version License: [Creative Commons CC BY 4.0 license](https://creativecommons.org/licenses/by/4.0/) Downloaded from: <https://hdl.handle.net/1887/135571>

Note: To cite this publication please use the final published version (if applicable).

SCIENTIFIC REPORTS

natureresearch

Received: 7 January 2019 Accepted: 23 July 2019 Published online: 02 August 2019

Efect of favor on neuronal OPENresponses of the hypothalamus and ventral tegmental area

A. M. vanOpstal1, A.A. van den Berg-Huysmans1, M. Hoeksma2, C. Blonk2, H. Pijl3, S.A. R. B. Rombouts1,4,5 & J. van derGrond1

Although it is well known that food intake is afected by the palatability of food, the actual efect of favoring on regulation of energy-homeostasis and reward perception by the brain, remains unclear. We investigated the efect of ethyl-butyrate (EB), a common non-caloric food favoring, on the blood oxygen level dependent (BOLD) response in the hypothalamus (important in regulating energy homeostasis) and ventral tegmental area (VTA; important in reward processes). The 16 study participants (18–25 years, BMI 20–23kg/m2) drank four study stimuli on separate visits using a crossover design during an fMRI setup in a randomized order. The stimuli were; plain water, water with EB, glucose solution (50gram/300ml) and glucose solution with EB. BOLD responses to ingestion of the stimuli were determined in the hypothalamus and VTA as a measure of changes in neuronal activity after ingestion. In the hypothalamus and VTA, glucose had a signifcant efect on the BOLD response but EB favoring did not. Glucose with and without EB led to similar decrease in hypothalamic BOLD response and glucose with EB resulted in a decrease in VTA BOLD response. Our results suggest that the changes in neuronal activity in the hypothalamus are mainly driven by energy ingestion and EB does not infuence the hypothalamic response. Signifcant changes in VTA neuronal activity are elicited by energy combined with favor.

In understanding the complexity of eating behavior, understanding the regulation of energy balance by is essential. Energy balance and intake is regulated by the brain by two major regulatory systems: the homeostatic system and the reward systems¹⁻³. The hypothalamus is known to control energy homeostasis through glucose and energy sensing and appetite regulation^{[1,](#page-7-0)[3](#page-7-1)}. The hypothalamus has been shown to respond directly to the ingestion of glucose and plays a pivotal role in central glucose sensing⁴⁻⁹. This central glucose sensing and metabolism have been established as an essential part of control of feeding and hunger¹⁰. In addition to the hypothalamus, hunger and appetite are also controlled in conjunction with the orbitofrontal cortex, insula and the reward system^{[1,](#page-7-0)11}. The reward, or mesolimbic, system is responsible for the hedonic response to food. An important brain area involved in this hedonic response is the ventral tegmental area (VTA). The VTA forms the basis of dopamine signaling in the mesolimbic system, which is a key substrate for reward prediction and response¹². Furthermore, the VTA is anatomically and functionally connected with the hypothalamus and integrates homeostatic signals with reward responses and therefore plays a pivotal role in regulating palatable feedin[g13–](#page-8-4)[15.](#page-8-5) Indeed, the VTA has been shown to have a (reward) response to both taste stimuli and the ingestion of energy $8,9,16$ $8,9,16$ $8,9,16$ $8,9,16$.

Pleasantness of food, afected by factors such as favor and texture, has a direct infuence on feeding behavior by making food palatable and attractive and thereby eliciting a reward response in the brain^{11,[13,](#page-8-4)17}. In this respect, pleasant flavoring might cause overeating and cause an energy misbalance¹⁸. Flavor of food might also be involved in energy balance as favor is an important part of the palatability of food and plays a role in satisfying and the rewarding effects and consumption volume of food^{17,19}. In this respect, pleasantness of food enhances the satiat-ing effect of high-fat and high-carbohydrate meals^{[20](#page-8-11)}. Additionally, flavor of food without ingestion of energy has been shown elicit responses from the brain and the periphery. Ingestion of a fat aroma with low fat content has been shown to elicit and modulate responses from the gustatory system²¹. Also, sweetness that is rated as pleasant

¹Leiden University Medical Center, Department of Radiology, Leiden, The Netherlands. ²Unilever Research & Development, Vlaardingen, The Netherlands. ³Leiden University Medical Center, Department of Internal Medicine, Section Endocrinology, Leiden, The Netherlands. ⁴Leiden Institute for Brain and Cognition (LIBC), Leiden, The Netherlands. ⁵Institute of Psychology, Leiden University, Leiden, The Netherlands. Correspondence and requests for materials should be addressed to A.M.O. (email: a.m.van_opstal@lumc.nl)

Table 1. Subject characteristics. Values in mean \pm standard deviation. Glucose and insulin levels in median and range. Glucose levels in mmol/L, normal fasted range: 3.9–5.5mmol/L, Insulin levels in mU/L, normal fasted range:<20 mU/L. *Average fasted blood levels over all four visits and the average post ingestion levels for two visits with glucose stimuli per subject measured 30minutes afer ingestion.

Table 2. Visual analogue scale (VAS) scores for hunger and flavor. Values in mean±standard deviation. VAS consisted of a 10 cm line scored from 0 to 10, anchors for the VAS score for hunger 0: 'not hungry' and 10: 'extremely hungry', anchors for the VAS score for favor 0: 'very unpleasant' and 10: 'very pleasant'

without caloric intake has been shown to elicit an insulin response when applied only to oral cavity^{22,23}. Although the evidence for the efect of taste on the cephalic phase insulin response is limited, these results do suggest that the favor of food, independent of energy content, might infuence the central regulation of the peripheral energy metabolism in addition to activation of the reward pathway in the brain. Furthermore, when glucose is paired with a congruent flavoring the perception of flavor is enhanced, and results in a greater feeling of satiety $24-26$. Therefore flavoring may have an effect on the responses in both the VTA and hypothalamus to glucose ingestion and energy ingestion in general. Additionally, the added perceived sweetness and satiation by favoring is interesting to investigate as this might be used as a strategy towards reduction of energy content as less nutritive sweetener could be used to reach the same level of overall sweetness^{[26](#page-8-16)}.

Taken together, earlier research indicates that favor may afect both homeostatic and hedonic aspects involved in energy intake. We aimed to gain a better understanding of the homeostatic and hedonic brain responses to favoring and energy content separately and the combination of both stimuli. Therefore we investigated the changes in neuronal activity, as measured by BOLD response, to the ingestion of glucose, Ethyl-Butyrate (EB) a common fruity food favoring, and a combination of both glucose and EB in the hypothalamus and VTA in normal weight young men. We hypothesized that favoring with EB, with and without glucose, could afect the neuronal activity in the hypothalamus and VTA.

Results

Subject characteristics. Sixteen participants were enrolled in the study. All participants successfully completed the all study visits. Table [1](#page-2-0) shows the subject characteristics of the study participants. All showed normal fasting glucose (3.9–5.5mmol/L) and insulin levels (<20 mIU/L) and these levels were not diferent between visits.

Hunger and favor scores. Subjective VAS scores for feelings of hunger and for rating of favor of the study stimuli are shown in Table [2.](#page-2-1) Before consumption, no significant differences were found between visits. The plain water and water flavored with EB stimuli initiated a significant increase in VAS scores for hunger. The flavor VAS score for water with EB was significantly lower compared with the other three study stimuli. The VAS score for favor for the combined stimulus of glucose and EB was not signifcantly diferent from water and glucose stimuli but was rated signifcantly more pleasant than EB alone.

Hypothalamic activity changes. Mean hypothalamic BOLD responses per study stimulus are shown in Fig. [1](#page-3-0) for illustration (top panel). Compared to baseline BOLD signal(before ingestion), ingestion of water led to a mean positive 0.63% BOLD response. Ingestion of water favored with EB led to a mean positive BOLD response of 0.18%. Both water with glucose and water with glucose combined with EB led to a mean negative BOLD response of −0.54% and −0.38%, respectively.

When determining the efect of energy content and the presence of EB favoring on the hypothalamic BOLD response we found that energy content had a significant effect on hypothalamic activity ($p=0.006$), however the EB flavoring did not have a significant effect ($p=0.430$). Energy content and EB flavoring did not have a significant $(p=0.310)$ interactive effect on the BOLD response, possibly due to the lack of effect of EB alone.

Figure 1. Hypothalamic mean BOLD signal change and BOLD time courses. BOLD responses to all four study stimuli (Mean % change with SEM). Percentage change from pre-ingestion (0–5minutes) to post-ingestion (minute 9–21 post ingestion) were calculated. P-values in the top panel for the diferences in percentage change between study stimuli were tested against the water intervention with linear mixed model analysis using the entire response. P-values in the bottom panel for diferences in percentage change between study stimuli were tested against the water intervention with linear mixed model analysis per time segment (min 9–13, min 14–17 and min 18–21). Grey areas in the time course graphs indicate period during which the stimuli were consumed, data points were excluded from the analysis.

Compared to the plain water response, water favored with EB did not lead to signifcant BOLD response $(p=0.204)$. On the contrary, both ingestion of glucose with and without EB resulted in a significant negative BOLD response (glucose only $p=0.008$, and glucose flavored with EB $p=0.017$). The addition of EB to glucose did not lead to significantly additive effect compared to glucose without flavoring ($p=0.870$). Ingestion of glucose flavored with EB ingested at 0 °C did not lead to different response compared to ingestion at room temperature. When looking at the response over diferent time periods (Fig. [1](#page-3-0) bottom panel, min 9–12, min 13–17 and min 18–21) we found that the response to glucose and glucose combined with EB was signifcantly diferent from water over all three time periods ($p=0.011$, $p=0.013$ and $p=0.023$ for glucose and $p=0.026$, $p=0.042$ and $p=0.025$ for glucose combined with EB).

Ventral tegmental area activity changes. Mean VTA BOLD responses per study stimulus are shown in Fig. [2](#page-4-0) for illustration (top panel). Compared to baseline BOLD signal (before ingestion), ingestion of water led to a mean positive 0.38% BOLD response. Ingestion of water favored with EB led to a mean positive BOLD response of 0.59%. Similar to the hypothalamic response, both water with glucose and water with glucose combined with EB led to a mean negative BOLD response of −0.31% and −0.93%.

When determining the efect of energy content and the presence of EB favoring on the overall BOLD response we found that energy content had a significant effect on VTA activity ($p=0.007$), however the EB flavoring did not have a significant effect ($p=0.612$). Energy content and EB flavoring did not have a significant ($p=0.363$) interactive efect on the BOLD response, possibly due to the lack of efect of EB alone.

Figure 2. Ventral Tegmental Area (VTA) mean BOLD signal change and BOLD time courses. BOLD responses to all four study stimuli (Mean % change with SEM). Percentage change from pre-ingestion (0–5minutes) to post-ingestion (minute 9–21 post ingestion) were calculated. P-values in the top panel for the diferences in percentage change between study stimuli were tested against the water intervention with linear mixed model analysis using the entire response. P-values in the bottom panel for diferences in percentage change between study stimuli were tested against the water intervention with linear mixed model analysis per time segment (min 9–13, min 14–17 and min 18–21). Grey areas in the time course graphs indicate period during which the stimuli were consumed, data points were excluded from the analysis.

Compared to the plain water response, water favored with EB did not lead to an altered BOLD response $(p=0.786)$. Ingestion of glucose only, resulted in a non-significant decrease in BOLD response ($p=0.187$). On the contrary, glucose flavored with EB mean difference led to a significant decrease in BOLD response ($p=0.028$), although this BOLD response was not significantly stronger compared to glucose only ($p=0.317$). Ingestion of glucose favored with EB ingested at 0 °C did not lead to diferent response compared to ingestion at room temperature. When looking at the response over diferent time periods (Fig. [2](#page-4-0) bottom panel, min 9–12, min 13–17 and min 18–21) we found that the response to glucose combined with EB was signifcantly diferent from water during the first minutes of the response ($p = 0.021$) but that this difference disappeared during the later minutes.

Associations between blood values, subjective ratings of hunger and favor and the hypothalamic and VTA activity changes. Mixed model analysis showed that increased ratings of hunger before stimulus ingestion were associated with a positive shif of the BOLD response in both the hypothalamus (+0.2%, $p=0.043$) and VTA (+0.2%, $p=0.012$). Moreover, ratings of hunger after stimulus ingestion were also associated with a positive shift of the BOLD response in both the hypothalamus $(+0.2\%, p=0.042)$ and VTA $(+0.03\%$, $p = 0.001$). Subjective rating of pleasantness of flavor did not have a significant association with the BOLD response in either the hypothalamus ($p=0.512$) and VTA ($p=0.573$). Blood glucose levels were not significantly associated with the BOLD response of the hypothalamus or VTA. Insulin levels afer ingestion were positively associated with the BOLD response in the VTA $(+0.2\% , p = 0.009)$.

Discussion

The results of this study show that in the VTA the combination of glucose with EB flavoring led to a significant decrease in neuronal activity, which was threefold stronger than to glucose alone. EB favoring in plain water (no caloric content) did not have any efect on the response of the VTA. In the hypothalamus the ingestion of glucose, and not EB, was associated with changes in neuronal activity.

The perception of sweetness, with and without energy content, has been shown to elicit a response from the reward system[13,](#page-8-4)[16.](#page-8-7) Sweetness is regarded by the brain as a predictor of energy content of the ingested food or drink^{16[,27](#page-8-17)[,28](#page-8-18)}. VTA is active in expectation of reward^{[13](#page-8-4)}, sweet flavoring could therefore lead to an increase in VTA activity because of the expectation of incoming energy content. Interesting in this context are non-nutritive sweeteners, which are increasingly used in foods and beverages, because they deliver sweet taste without any caloric content^{29,30}. However, the flavoring we used in our study is not necessarily perceived as sweet on its own without the addition of glucose or another sweetener. Additionally, the EB stimulus was rated as less pleasant then the other stimuli and therefore might not have been rewarding by itself. Tis could explain why we did not fnd a signifcant response from the VTA, although the activity did appear to increase, with just the EB favoring added to plain water^{[8](#page-8-6),[9,](#page-8-0)[16](#page-8-7)}. Additionally, EB could have effects on other brain areas that we missed by measuring only the responses of the hypothalamus and VTA.

Adding favoring to sugars has been shown to amplify the sweet taste and palatability which could increase the rewarding response^{[19](#page-8-10),[20](#page-8-11)}. Additionally, when glucose is paired with a congruent flavoring, the perception of flavor is also enhanced^{[24,](#page-8-15)25}, and could subsequently lead to a stronger reward response. This is in line with our finding that EB combined with glucose elicited the most pronounced response from the VTA. Furthermore, energy coupled with favoring has been shown to lead to stronger subjective feelings of satiation and short term satiety compared to either stimulus separately²⁶, which can also be linked to a stronger reward response by the brain. The strongest response to the combined stimulus in the VTA could be explained by the VTAs role in regulating palatable eating by integrating energy content with reward via the VTAs anatomical and functional connections with the hypothalamus^{13–[15](#page-8-5)}. The strongest response to the combined stimulus could also be explained by the fact that this stimulus could be the most intense perceptual stimulation that afects multiple sensory modalities including both gustatory and olfactory aspects but also trigeminal sensation. However, we did not measure perceptual intensity ratings of the stimuli so we cannot determine of this was the case. The strongest effect of combined flavoring and glucose on the VTA could indicate that added favoring can be used as strategy towards reduction of energy content, as our results indicate that the response to energy content could be strengthened when combined with favoring therefore allowing for a possible lower total energy content.

In addition to reward efects, favor might afect the neuronal activity of the hypothalamus as sweet taste without caloric intake has been shown to elicit an insulin response^{[22](#page-8-13)}. However, other studies indicate that taste only elicits or modulates a hormonal response in combination with caloric content, as regulated by the hypothal-amus^{[5](#page-7-3)[,30](#page-8-20)[,31](#page-8-22)}. Earlier studies by our group have shown that the ingestion of glucose solution leads to a decreased activity throughout the brain and specifically in the hypothalamus^{8,[9](#page-8-0),[32](#page-8-23)}. The results from the current study confirm once again that hypothalamic activity decreases afer glucose ingestion and show that the driving force of the hypothalamic response is indeed the ingestion of energy and not the favor indicated by the lack of response to favoring without energy content. Our result are also concordant with the results of other previous studies show-ing that flavor without energy does not affect the hypothalamus^{4-[7](#page-8-24)}.

Our data show that stronger feelings of hunger, both before and afer stimulus ingestion, had a positive efect on activity change adjusted for treatment in both the hypothalamus and VTA. Tis indicates that a stronger feeling of hunger lessened the decrease in activity in the hypothalamus. In a state of hunger, the brain - and specif-cally the hypothalamus and VTA - have been shown to have a high activity^{[5,](#page-7-3)[6,](#page-8-25)[13,](#page-8-4)33}, which could thus decrease after energy intake and with satiety. Our fnding that a higher subjective score for hunger leads to a larger decrease in activity in the hypothalamus and the VTA support this theory. This suggests that these responses might reflect an objective evaluation of (perceived) energy content and reward.

A limitation of this study is the generalizability as we only investigated a relatively small sample of male volunteers and it can be expected that sex diferences are present, since it is known that there are several sex-specifc differences in energy metabolis[m34.](#page-8-27) Additionally, we only investigated lean subjects and earlier studies have shown different hypothalamic function in obesity³⁵. A limitation of our study design was the use of water as a negative control. Water might not be a negative control as could elicit a response a response from the hypothalamus as fuid balance is also regulated by the hypothalamus. However, since all our stimuli contained or were dissolved in water we this efect would be found with all stimuli. We did not measure thirst in our participants and therefore we could not correct for the efect of this per study visit on a possible response of the hypothalamus to regulate fuid balance. Furthermore, our combined stimulus of EB with glucose is a very complicated stimulus with gustatory and olfactory effects. The combined stimulus could lead to various combined effects on that might be different from just the additive efects of glucose plus favoring. Unfortunately our current design does not allow us the determine these efects. A strength of our study was our crossover study design, which allowed for a reliable within-subject comparison between interventions as participants were their own controls.

Taken together, our results suggest that the response of the VTA is strongest to a combination of both favor and energy and that the hypothalamic response is mainly driven by energy ingestion. Although neither response seemed dependent on subjective ratings of pleasantness in the basal brain areas investigated here, the levels of hunger do influence the responses in these areas. This suggests that the responses by the hypothalamus and VTA might be mostly involved in subconscious rather the conscious regulation of satiety, appetite and feeding behavior and is mainly driven by energy demand and intake.

Figure 3. Study procedures. Illustrative depiction of the study procedures and timing during the study visits.

Methods

Study participants. Sixteen healthy, normal-weight men participated in our study. Participants were recruited via local advertisements and through the use of mailing lists. All participants were between the ages of 18 and 25 and had a body mass index (BMI) ranging from 20 to 23. Exclusion criteria were the following: a history of diabetes or disturbed glucose metabolism; any genetic, psychiatric, renal, hepatic or chronic disease; recent fluctuations in weight >3 kg; current smoking; current alcohol consumption >21 units/week and use of recreational drugs; recent blood donation or participation in other biomedical research (within the last 3 months); use of medication affecting glucose of lipid metabolism; contra-indications for MRI-scanning. The protocol was approved by the Medical Ethics Committee of the Leiden University Medical Center (LUMC) and registered at clinicaltrails.gov under number NCT03202342. All investigations have been conducted according to the principles expressed in the Declaration of Helsinki. Written informed consent was obtained from all participants afer complete written and verbal description of the study was given.

Experimental procedure. We used a randomized, controlled, crossover study design. A batch-wise randomization procedure, respecting 1st order, was followed. The randomization was performed by an independent person. Participants arrived at MRI facilities of the Leiden University Medical Centre (LUMC) afer an overnight fast on a separate study occasion for each stimulus. Afer blood sampling and recording pre scanning visual analogue scores (VAS), participants were positioned in the MRI scanner where the study stimuli were ingested within the scanner bore through a per oral tube during fMRI scanning. Study procedures are illustrated in Fig. [3.](#page-6-0) The different stimuli used and investigated were; plain water, water with ethyl-butyrate (EB), glucose solution and glucose solution with ethyl-butyrate (EB) (glucose solutions consisted of 50 grams of glucose dissolved in 300ml water),stimuli were ingested at room temperature. Water was plain, non-chlorinated tap water. Ethyl-butyrate is a commonly used food favoring with a fruity favor and is an FDA and EU approved food favoring under title 21 section 182.60 and EU Regulation 1334/2008 & 178/2002. EB was delivered in a liquid form dissolved in glycerol by Unilever Research and Development Vlaardingen B.V. EB and was frst diluted to a lower concentration and then added to the 300 ml plain water or glucose solution to a concentration of 0.01%. The stimuli were tasted before the start of the study to determine the dose of EB that would be appropriate. Glucose samples were perceived to be sweetest. EB alone dissolved in water was perceived as slightly sweeter than water, but when combined with glucose did not enhance the sweetness when compared to glucose alone. EB alone as well as glucose alone were perceived to be fruity, and the combination of glucose with EB resulted in an enhancement of fruitiness. In the present trial, an additional ffh stimulus was investigated, consisting of a glucose solution with EB ingested at 0 °C, to investigate the efects of temperature. As the aim of the current study was to determine the effects of flavoring only and the effects of temperature have been described in an earlier study by our group^{[8](#page-8-6)} the results of this condition are therefore only briefly shown and not discussed further in the current study. The stimuli were ingested fve minutes afer the start of the fMRI scan.

Blood sampling and VAS scores. Blood samples were used to ascertain a normal glucose metabolism in each participant. On every test day, 2 blood samples (5ml each) were drawn by venipuncture in an antecubital vein; one sample was taken before scanning and the other one afer the scanning procedure, 30 minutes afer ingestion of the study solution. Sample handling and analysis was performed by the laboratory for Clinical Chemistry at LUMC. Plasma glucose was measured using a fully automated Hitachi 704/911 system. Plasma insulin was measured by a Radio Immuno Assay (Medgenix, Fleurus, Belgium). A total amount of 10ml blood was taken on each study day. Participants were asked to rate their feelings of hunger in advance of the scanning procedure and aferwards, using a visual analogue scale (VAS) which consisted of a 10cm line, with 'not hungry' and 'extremely hungry' as anchors. Additionally, afer the MRI scan pleasantness of the drink was rated using a similar VAS using 'very unpleasant' to 'very pleasant' as anchors.

MRI data acquisition. The MRI was performed using a 3 Tesla whole body MRI scanner (Philips Achieva, Philips Healthcare, Best, The Netherlands) equipped with a 32-channel head coil. The protocol for structural MRI comprised a scout view for planning, a high resolution 3DT1-weighted sequence for registration purposes and a mid-sagittal high resolution single slice for accurate hypothalamus and VTA localization (repetition time 550ms, echo time 10 ms, field of view 208×208 mm, voxel size = $0.52 \times 0.52 \times 14$ mm). Mid-sagittal fMRI was performed for 21 minutes in total, by a T_2^* weighted, gradient echo-planar imaging (EPI) sequence mid-sagittal single slice that renders BOLD contrast (repetition time 120 ms; echo time 30 ms; flip angle 30 \degree ; field of view 208 \times 208 mm; voxel size = $0.81 \times 0.90 \times 14$ mm; 21 k-line excitations for each dynamic volume, scanning time of one dynamic image 2.52 seconds for a total of 500 data points). A slice thickness of 14mm was chosen to encompass the hypothalamus in the left to right direction and a single slice technique was used for sufficient signal to noise ratio.

High resolution mid brain functional MRI analysis. Data was pre-processed as described previously^{[7](#page-8-24)}. In short data was averaged for each set of 4 subsequent volumes, reducing the 500 dynamic scans to 125 time

Figure 4. A representative sagittal view with the hypothalamus (A) and the VTA (B) ROIs. For (A), the optic chiasm (1), the mammillary bodies (2), the thalamus (3) and the anterior commissure (4) were used as landmarks. For (B), half the volume of (A), the top of the cerebral aqueduct (5) and the mammillary bodies (2) were used as landmarks. The reference ROI was drawn superior of the genu to the corpus callosum (6) in the grey matter.

points used for further analysis. Regions of interests (ROIs) were drawn manually, using subject-specific T_1 images to define anatomical borders. Three ROIs were drawn: the hypothalamus, the VTA and a reference area. The ROI for the hypothalamus was drawn as described earlier using the optic chiasm, mammillary bodies, thal-amus and anterior commissure as anatomical landmarks^{[7](#page-8-24)}. Using literature describing the VTA region^{36,[37](#page-8-30)} we defned the ROI for the VTA in the midbrain using the top of the cerebral aqueduct and the mammillary bodies as anatomical landmarks (example ROIs shown in Fig. [4\)](#page-7-4). To correct for scanner drif, a third internal reference ROI was selected superior to the genu of the corpus callosum in the grey matter. BOLD signals derived from the hypothalamus and the VTA were corrected for BOLD signals obtained from this reference ROI by dividing the ROI BOLD signals by the reference BOLD signal. Percentage BOLD change from corrected baseline was calculated by dividing the post ingestion BOLD signal by the average baseline BOLD signal for the individual post ingestion dynamic scans/measurement time points ($n=125$) and for the total post ingestion period (minute 8–21 of the fMRI scan).

Statistical analysis. Statistical analysis was performed using SPSS version 23. Diferences in blood levels and VAS scores between pre- and post-ingestion and between study stimuli were tested with repeated measure ANOVA. All fMRI results are reported as BOLD change relative to baseline (0–5 minutes pre-drinking BOLD signal). To determine the main effect of energy content (presence of glucose yes or no) and flavor (presence of EB yes or no) on the study stimuli linear mixed model analysis was performed using energy content and the presence of favoring and the study occasion as fxed efects and subject*study occasion as a random factor and the BOLD measurement time points as a covariate. To determine the interaction between energy content and favoring the same mixed model was performed with an interaction term of both factors as a fxed efect. Statistical analysis of the diference in treatment efect between the study stimuli was performed by similar linear mixed model analysis using the study stimulus and study occasion as fxed efects, time point as a covariate and subject*occasion as a random factor using plain water as a negative and glucose solution as a positive reference stimulus. To determine the association between blood levels and the subjective ratings of hunger and favor of the study stimulus and the BOLD response of the hypothalamus and VTA additional similar mixed models were used where these variables were added as covariates. All these mixed model analyses were applied on the n=125 point dataset and therefore consider the entire response. To determine whether the diference in treatment efect changed during the measured time period we performed additional mixed model analysis over three time bins (min 9–12, min 13–17 and min 18–21).

Data Availability

The raw data supporting the conclusions of this manuscript will be made available by the authors, without undue reservation, to any qualifed researcher.

References

- 1. Morton, G. J. *et al*. Central nervous system control of food intake and body weight. *Nature* **443**(7109), 289–295 (2006).
- 2. Morton, G. J., Meek, T. H. & Schwartz, M. W. Neurobiology of food intake in health and disease. *Nat. Rev. Neurosci* **15**(6), 367–378 (2014).
- 3. Schwartz, M. W. *et al*. Central nervous system control of food intake. *Nature* **404**(6778), 661–671 (2000).
- 4. Smeets, P. A. *et al*. Functional magnetic resonance imaging of human hypothalamic responses to sweet taste and calories. *Am. J. Clin. Nutr* **82**(5), 1011–1016 (2005).
- 5. Smeets, P. A. *et al*. Functional MRI of human hypothalamic responses following glucose ingestion. *Neuroimage* **24**(2), 363–368 (2005).
- 6. Smeets, P. A. *et al*. Oral glucose intake inhibits hypothalamic neuronal activity more efectively than glucose infusion. *Am. J. Physiol Endocrinol. Metab* **293**(3), E754–E758 (2007).
- 7. Teeuwisse, W. M. *et al*. Short-term caloric restriction normalizes hypothalamic neuronal responsiveness to glucose ingestion in patients with type 2 diabetes. *Diabetes* **61**(12), 3255–3259 (2012).
- 8. van Opstal, A. M. *et al.* The effect of consumption temperature on the homeostatic and hedonic responses to glucose ingestion in the hypothalamus and the reward system. *Am. J. Clin. Nutr* **107**(1), 20–25 (2018).
- 9. van Opstal, A. M. *et al*. Dietary sugars and non-caloric sweeteners elicit diferent homeostatic and hedonic responses in the brain. *Nutrition*, 2018.
- 10. Davis, J. D. *et al*. Sustained intracerebroventricular infusion of brain fuels reduces body weight and food intake in rats. *Science* **212**(4490), 81–83 (1981).
- 11. Berthoud, H. R. & Morrison, C. The brain, appetite, and obesity. Annu. Rev. Psychol 59, 55-92 (2008).
- 12. Schultz, W., Dayan, P. & Montague, P. R. A neural substrate of prediction and reward. *Science* **275**(5306), 1593–1599 (1997).
- 13. Meye, F. J. & Adan, R. A. Feelings about food: the ventral tegmental area in food reward and emotional eating. *Trends Pharmacol. Sci* **35**(1), 31–40 (2014).
- 14. Nieh, E. H. *et al*. Inhibitory Input from the Lateral Hypothalamus to the Ventral Tegmental Area Disinhibits Dopamine Neurons and Promotes Behavioral Activation. *Neuron* **90**(6), 1286–1298 (2016).
- 15. Sheng, Z. *et al*. Metabolic regulation of lateral hypothalamic glucose-inhibited orexin neurons may infuence midbrain reward neurocircuitry. *Mol. Cell Neurosci* **62**, 30–41 (2014).
- 16. Haase, L., Cerf-Ducastel, B. & Murphy, C. Cortical activation in response to pure taste stimuli during the physiological states of hunger and satiety. *Neuroimage* **44**(3), 1008–1021 (2009).
- 17. Bellisle, F. *et al*. Sweetness, satiation, and satiety. *J. Nutr* **142**(6), 1149S–1154S (2012).
- 18. Szalay, C. *et al*. Gustatory perception alterations in obesity: an fMRI study. *Brain Res* **1473**, 131–140 (2012).
- 19. Rolls, E. T. Taste, olfactory, and food texture processing in the brain, and the control of food intake. *Physiol Behav* **85**(1), 45–56 (2005). 20. Warwick, Z. S. *et al*. Taste and smell sensations enhance the satiating efect of both a high-carbohydrate and a high-fat meal in
- humans. *Physiol Behav* **53**(3), 553–563 (1993). 21. Frank, S. *et al*. Olive oil aroma extract modulates cerebral blood fow in gustatory brain areas in humans. *Am J Clin Nutr* **98**(5), 1360–6 (2013).
- 22. Just, T. *et al*. Cephalic phase insulin release in healthy humans afer taste stimulation? *Appetite* **51**(3), 622–627 (2008).
- 23. Abdallah, L., Chabert, M. & Louis-Sylvestre, J. Cephalic phase responses to sweet taste. *Am. J. Clin. Nutr* **65**(3), 737–743 (1997).
- 24. Frank, A. R. & Byram, J. Taste-smell interactions are tastant and odorant dependent. *Chem. Senses* **13**, 445–455 (1988).
- 25. Green, B. G. *et al*. Enhancement of retronasal odors by taste. *Chem. Senses* **37**(1), 77–86 (2012).
- 26. Yin, W. *et al*. Efects of aroma and taste, independently or in combination, on appetite sensation and subsequent food intake. *Appetite* **114**, 265–274 (2017).
- 27. Avena, N. M., Rada, P. & Hoebel, B. G. Evidence for sugar addiction: behavioral and neurochemical efects of intermittent, excessive sugar intake. *Neurosci. Biobehav. Rev* **32**(1), 20–39 (2008).
- 28. Swithers, S. E. & Davidson, T. L. A role for sweet taste: calorie predictive relations in energy regulation by rats. *Behav. Neurosci* **122**(1), 161–173 (2008).
- 29. Brown, R. J., de Banate, M. A. & Rother, K. I. Artifcial sweeteners: a systematic review of metabolic efects in youth. *Int. J. Pediatr. Obes* **5**(4), 305–312 (2010).
- 30. Brown, R. J. & Rother, K. I. Non-nutritive sweeteners and their role in the gastrointestinal tract. *J. Clin. Endocrinol. Metab* **97**(8), 2597–2605 (2012).
- 31. Renwick, A. G. & Molinary, S. V. Sweet-taste receptors, low-energy sweeteners, glucose absorption and insulin release. *Br. J. Nutr* **104**(10), 1415–1420 (2010).
- 32. van Opstal, A. M. *et al*. Brain activity and connectivity changes in response to glucose ingestion. Nutr Neurosci, p. 1–8 (2018).
- 33. Trivedi, B. P. Neuroscience: Dissecting appetite. *Nature* **508**(7496), S64–S65 (2014).
- 34. Varlamov, O., Bethea, C. L. & Roberts, C. T. Jr. Sex-specifc diferences in lipid and glucose metabolism. *Front Endocrinol. (Lausanne)* **5**, 241 (2014).
- 35. Matsuda, M. *et al*. Altered hypothalamic function in response to glucose ingestion in obese humans. *Diabetes* **48**(9), 1801–1806 (1999).
- 36. D'Ardenne, K. *et al*. BOLD responses refecting dopaminergic signals in the human ventral tegmental area. *Science* **319**(5867), 1264–1267 (2008).
- 37. Krebs, R. M. *et al*. Novelty increases the mesolimbic functional connectivity of the substantia nigra/ventral tegmental area (SN/ VTA) during reward anticipation: Evidence from high-resolution fMRI. *Neuroimage* **58**(2), 647–655 (2011).

Author Contributions

J.G., A.B., M.H., H.P. and C.B. designed research; A.O. and J.G. conducted research; A.O. and A.B. analyzed data; A.O., J.G. and S.R. wrote the paper; J.G. and S.R. had primary responsibility for fnal content. All authors read and approved the fnal manuscript.

Additional Information

Competing Interests: Tis study was funded by Unilever Research and Development Vlaardingen B.V. The Netherlands. M. Hoeksma and C. Blonk are both employees of Unilever Research and Development Vlaardingen B.V. The Netherlands. All other authors have no competing interests to disclose.

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional afliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit [http://creativecommons.org/licenses/by/4.0/.](http://creativecommons.org/licenses/by/4.0/)

 \circ The Author(s) 2019