Neural dissociation of food and money related reward processing using an abstract incentive delay task

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ABSTRACT

Food is an innate reward stimulus related to energy homeostasis and survival, whereas money is considered a more general reward stimulus that gains a rewarding value through learning experiences. Although the underlying neural processing for both modalities of reward has been investigated independently from one another, a more detailed investigation of neural similarities and/or differences between food and monetary reward is still missing. Here, we investigated the neural processing of food compared to monetary related rewards in 27 healthy, normal-weight women using functional magnetic resonance imaging (fMRI). We developed a task distinguishing between the anticipation and the receipt of either abstract food or monetary reward. Both tasks activated the ventral striatum during the expectation of a reward. Compared to money, greater food related activations were observed in prefrontal, parietal and central midline structures during the anticipation and lateral orbitofrontal cortex (lOFC) during the receipt of food reward. Furthermore, during the receipt of food reward, brain activation in the secondary taste cortex was positively related to the BMI. These results indicate that food dependent activations encompass to a greater extent brain regions involved in self-control and self-reflection during the anticipation and phylogenetically older parts of the lOFC during the receipt of reward.

Keywords: neural processing of food rewards, eating disorders, BMI, ventral striatum, orbitofrontal cortex.
INTRODUCTION

Motivation for food intake in humans is not only dependent on energy homeostasis but is also determined by hedonic aspects of food intake. From an evolutionary account, the latter constitute important factors formerly ensuring survival in times when food supply was rare. However, in our present obesogenic environment, a heightened food reward sensitivity facilitates the overconsumption of food. As a result, the obesity epidemic has become a serious public health and socioeconomic problem (Swinburn et al., 2011). In recent years, brain imaging studies have shown that alterations in brain reward pathways influence food intake (Stice et al., 2009). These findings offer promising routes to identify neural markers that may underlie the overconsumption of food and other pathological eating behaviors (Morton et al., 2006).

Anticipated intake of high-calorie food leads to an activation of the mesocorticolimibic reward system (O’Doherty et al., 2002; Pelchat et al., 2004). In contrast, consummatory food reward mainly involves the primary (insula/ frontal operculum), as well as the secondary gustatory cortex (lateral orbitofrontal cortex (lOFC)) (O’Doherty et al., 2002; Rolls, 2007). Nevertheless, activations in these regions are not only restricted to the consumption of palatable food but can also be found during the exposure to appetizing visual food stimuli (Beaver et al., 2006; Porubská et al., 2006), suggesting that abstract visual stimuli can inherit similar motivational and hedonic qualities. Furthermore, previous studies have shown that direct and indirect measures of food intake such as the body mass index (BMI) or frequency of overeating, are directly related to activations in reward related brain regions such as the ventral striatum (VS) and OFC during the processing of food stimuli (Lawrence et al., 2012; Murdaugh et al., 2012; Stoeckel et al., 2008). However, the direction of this
association remains unclear as some studies report a positive relation between weight and activity in reward related regions, whereas other studies report the opposite (Burger and Stice, 2011; Ziauddeen et al., 2012).

Given the evolutionary relevance of food reward, one would expect that the processing of food as an innate reward stimulus differs to some extent from more general monetary reward processing. However, there is a paucity of studies directly comparing food and monetary reward (Metereau and Dreher, 2013; Sescousse et al., 2013). Previous studies have assessed food and secondary reward processing independently from one another. These studies found both common, as well as distinct activation patterns in the brain reward network for the two reward modalities (Diekhof et al., 2012; Sescousse et al., 2013). Specifically, learned reward associations (i.e. money) are additionally represented in evolutionary more recent brain regions, whereas innate food rewards also recruit the somatosensory cortex (Sescousse et al., 2013). In the current study, we were interested in assessing both the anticipation as well as the receipt of food and monetary rewards intra-individually. To our knowledge, this is the first study directly comparing food and monetary reward using the same experimental task for both modalities of reward.

Since the VS and medial OFC (mOFC) have been strongly related to subjective experiences of pleasantness (Kühn and Gallinat, 2012), we hypothesized that both the VS and mOFC will show significant activations independently from the reward modality (Knutson et al., 2001b; Morton et al., 2006). Additionally, we expected that food rewards will be associated with higher general activation during the anticipation phase and greater activation in gustatory brain regions during the receipt phase (frontal operculum and IOFC, respectively). Furthermore, we expected that the latter observation will be modulated by the individual participant's BMI.
SUBJECTS AND METHODS

Subjects

Thirty healthy right-handed female adults participated in the study. Mean age was 26.5 years (± 6.3), mean BMI was 21.8kg/m² (± 1.8). Data from two subjects was excluded from the analysis due to excessive head-movement (greater than 3 mm in any one direction); data from one subject was excluded due to a technical error, leaving 27 subjects for the final analysis. All had normal or corrected to normal vision, and were screened for medical and psychic diseases by taking the medical history, measuring body weight and height, and by interviewing them with the Structured Clinical Interview for DSM-IV (SCID, Wittchen, 1997). Chronically-ill, overweight and participants with a lifetime psychic diagnoses were excluded. Furthermore, participants were excluded if they were regular smokers. The present study complies with the Code of Ethics of the World Medical Association (Declaration of Helsinki, version 2008) and was approved by the Ethics Committee of the Medical School of the University of Heidelberg. Written informed consent was obtained from all participants after the procedures had been fully explained.

Study procedure

All participants were asked to come to the clinic without having breakfast and to refrain from consuming alcoholic drinks for 24 hours before the experiment. The study began with a light standardized breakfast at 9.00 a.m., followed by the SCID. After the interview, all subjects were asked to conduct a battery of neuropsychological tasks and to complete a package of questionnaires. The findings
of the latter are not reported in this study. The MRI scanning was performed for all subjects at 12:00 p.m., corresponding to the lunchtime of most of the participants.

Stimuli and task

We used a modified version of the “monetary incentive delay” (MID) task as proposed by Abler and colleagues (Abler et al., 2005; Knutson et al., 2001a). In a previous study, we showed that this paradigm allows an efficient probing of both anticipation and consumption of reward (Simon et al., 2010). Additionally, we employed a “food incentive delay” (FID) task: instead of money, subjects were able to win “snack points” (SP), which they then could exchange for sweet and salty snacks immediately after the MRI measurement (Simon et al., 2014). There were four blocks of reward tasks, consisting of 55 trials each. The block sequence was either SMSM (S=Snacks, M=Money), or MSMS, and was counterbalanced over the participants. Before entering the scanner, subjects performed a practice version of the task lasting three minutes for each condition for which they received neither payment nor snacks. The degree of potential rewards varied on three levels as indicated via graphical cues (see figure 1). In both tasks, each trial started with the presentation of a symbol (“cue”, 750 ms) indicating the amount of money/ number of snack-points they could win with a correct response (i.e., 1 EUR, 20 cents, 0 EUR, or 10 SP, 2 SP, 0SP, respectively). After an anticipation period (“delay”, 3.000 ms) subjects had to correctly react to one of two symbols (“targets”; i.e., triangle inclined to the right or a triangle inclined to the left) with a left or right button press corresponding to the direction of the triangle (index or middle finger of dominant hand) within a fixed interval of 1,000 ms. This leads to a low task-difficulty with a very high success rate, independently of the reaction speed of the participant. In order to guarantee a steady rate of reward vs. non-reward throughout all subjects, we used a probabilistic reward pattern, i.e.
reward was not paid out in 30 predefined trials (out of the 80 reward trials).

Immediately after target presentation, feedback appeared (“feedback”, 1.500 ms), notifying participants about the amount of money/ snack points they had won and about their cumulative total. In order to increase statistical efficiency, trials were separated by jittered intertrial intervals (ITIs) ranging from 1 to 8 s, with a mean of 3.5 s. The MID task used graphical depictions showing a wallet filled with the corresponding amount of money won during each trial. The FID task used pictures of either a large basket filled with snacks, a small basket filled with snacks or an empty basket, depending on the amount of snack points won. An incorrect button press resulted in zero payout. To ensure that subjects paid attention and responded to every experimental condition, a penalty of -1 EUR/ -10 SP was applied if they missed to press one of the two buttons. In the MID task, subjects were able to win a maximum of 30 EUR. In the FID task, the maximum amount to be won was 300 SP, with any snack of the basket being 50 SP worth.

fMRI acquisition

Images were collected using a 3-T Siemens Trio MRI scanner (Siemens Medical Solutions, Erlangen, Germany) equipped with a standard 32 channel head coil. The subjects performed four functional runs lasting 9.3 min each with 280 volumes per run. In order to minimize susceptibility artifacts in the OFC 30 oblique slices (interleaved acquisition) with a 10° angle relative to the AC-PC axis were acquired with 1 mm interslice gap, using a T2*-sensitive single-shot EPI sequence with following parameters: TR=2000 ms, TE=30 ms, resulting in an in-plane resolution of 3×3×4 mm³, flip angle=80°,field of view=192×192 mm. Participants viewed visual stimuli on a projection screen via a mirror fixed to the head coil and responded with the right hand using a button box. High-resolution T1 MPRAGE anatomical images
were acquired (192 slices, voxel size 1×1×1 mm, TR 1570 ms, TE 2.63 ms, 9° flip angle) for anatomical reference.

fMRI data analysis

Functional MRI data were pre-processed and analyzed with SPM8 (Statistical Parametric Mapping, Wellcome Department of Cognitive Neurology, London, UK). To account for magnetic field equilibration, four volumes from the start of each functional run were excluded from analysis. Pre-processing of functional scans included slice time correction (reference to the first slice, using SPM8's Fourier phase shift interpolation), within-subject registration and unwarping of time-series (to correct for motion artifacts), coregistration of the T1 image with the mean T2*-image, spatial normalization of both the functional and structural images to a standard T1 template brain (ICBM152, Montreal Neurological Institute, MNI), resulting in a voxel size of 3 mm\(^3\) for functional images and a voxel size of 1 mm\(^3\) for high resolution anatomic images and smoothing with an 8 mm full-width half-maximum isotropic Gaussian kernel. A 128-s high-pass filter was used to remove low-frequency noise and signal drift.

At the first level of analysis, pre-processed functional MRI data for both tasks were analyzed separately in the context of the general linear model (GLM) approach (Friston et al., 1994). Regressors were modeled separately for the three different anticipation phases (anticipation of 1 EUR, 20 cents, and 0 EUR for the MID task, anticipation of 10 SP, 2 SP, and 0 SP for the FID task) and the five different outcome phases (receipt of 1 EUR, omission of 1 EUR, receipt of 20 cents, omission of 20 cents, and receipt of 0 EUR/neutral outcome for the MID task, receipt of 10 SP, omission of 10 SP, receipt of 2 SP, omission of 2 SP, and receipt of 0 SP/neutral outcome for the FID task) as explanatory variables convolved with the gamma-variate
function described by Cohen (1997). Targets and error trials were included as additional regressors of no interest. Linear combinations of the estimated GLM parameters allow the assessment of changes in the BOLD responses of individual participants, contingent on the experimental condition. Individual contrast images corresponding to the effects of interest were subsequently constructed. For the analysis of reward anticipation, we contrasted the anticipation of a high reward (1 EUR or 10 SP) with the anticipation of no reward (0 EUR or 0 SP, \textit{anticipation\_high}). For the analysis of the impact of a rewarding outcome, we contrasted the receipt of a high reward (1 EUR or 10 SP) with the receipt of no reward (0 EUR or 0 SP, \textit{receipt\_high}), controlling for the anticipation phase which preceded both outcome types. All contrasts were modeled separately for each task.

At the second level of analysis, the individual contrast images of all participants were included in a random-effects analysis, allowing population inference (Holmes and Friston, 1998). A whole brain analysis using the specific contrasts of interests was carried out in order to identify reward-sensitive brain areas. Within-group activation was compared using a one-sample t-test, and between-tasks activation using a two-sample t-test. We report results significant at an uncorrected voxel level threshold of \(p<0.001\) (Spreckelmeyer et al., 2009). The location of the peak activity associated with each cluster of activation is reported in MNI-coordinates.

**Region of interest (ROI) analysis**

In order to assess brain activation in reward related brain regions, we constructed functional ROIs based on the results of the whole brain analysis during the FID task, using a threshold of \(p<0.001\) uncorrected. Functional ROIs were constructed based on activation clusters found in reward related regions during both the anticipation and the receipt of food rewards. Spheres of 8 mm diameter were created around the
respective peak voxels of activation. We constructed four ROIs: left VS, right VS, left
IOFC, and right IOFC. The left and right VS ROIs were based on peak activation in
the left (MNI-coordinates: x = -6, y = 5, z = 2) and right (MNI: 12, 2, 2) VS
respectively, passing a threshold of p<0.001 during the contrast anticipation_high.
The left (MNI: -39, 47, -6) and right (MNI: 42, 41, -14) IOFC ROIs were constructed
using peak activation in IOFC regions during the contrast receipt_high. Mean percent
signal change was extracted for each ROI using MarsBaR (Brett et al., 2002). Simple
correlation (Pearson’s r) analyses were performed for psychometric scales and mean
percent signal change in the ROIs using SPSS version 20.
RESULTS

Behavioral results

Reaction times and error rates for each task are given in table S1 in the supplementary materials. We did not observe a significant difference in reaction time or error rate between tasks ($p > .29$).

Neuroimaging results

Anticipation of reward

During the contrast $\text{anticipation\_high}$, the FID task displayed activations in the bilateral VS, bilateral thalamus, subgenual anterior cingulate cortex (ACC), right anterior insula and bilateral occipital cortex (table 1, figure 2). The cluster in the right VS extends to the anterior thalamus. For the MID task, we observed activations in the left VS, right anterior insula, bilateral thalamus and a portion of the midbrain (table 1). The extracted signal change at the VS ROI for the FID task during different reward conditions (anticipation of 10 SP, 2 SP, 0 SP) showed a noticeable linear decrease from 10 SP via 2 SP to 0 SP for both the lVS and rVS (see figure 2).

Receipt of reward

During the contrast $\text{receipt\_high}$, the FID task showed activations in the bilateral OFC, bilateral inferior IPFC, left inferior operculum, bilateral parietal cortex, and bilateral occipitotemporal cortex (table 1, figure 3). For the MID task, we observed a large cluster with punctum maximum in the subgenual ACC, extending into the mOFC. Additionally, we observed activity in the bilateral inferior IPFC, left anterior insula, posterior cingulum, bilateral parietal, and occipitotemporal cortex (table 1).
The extracted signal change showed a linear decrease from the receipt of 10 SP via 2 SP to 0 SP for both the left and right IOFC (see figure 3) in the FID task.

*Comparison between monetary and food reward*

Compared to the MID task, the FID task showed significant differences in brain activation using a whole brain analysis. During *anticipation_high*, the FID task showed stronger activations in the inferior IPFC, OFC, medial frontal cortex, posterior cingulum, angular gyrus, and middle temporal gyrus. During the *receipt_high*, the FID task showed stronger activations in the posterior IOFC and occipital cortex (table 1, figure 4). There were no brain regions that showed greater brain activations in the MID compared to the FID task.

*Correlational analysis*

We found a positive correlation for the FID task between percent signal change extracted from the right and left IOFC ROI during the *receipt_high* contrast and the BMI of the individual participants (figure 3). There were no significant correlations for the FID task between signal change extracted from the left and right VS and the BMI. For the MID task, no correlations between the signal change of the VS nor the IOFC, or the individual BMI were observed.
4. DISCUSSION

The aim of this study was to investigate common and distinct patterns in neural processing of food- and more general monetary related rewards in healthy individuals. In line with our hypotheses, we observed an increased activation in the VS during the anticipation of reward, irrespective of the reward modality. Contrary to our assumptions, we only observed mOFC activity during the receipt of monetary rewards, whereas the receipt of food reward was associated with heightened activation of the bilateral lOFC. Furthermore, we observed increased activation in the lateral ventral prefrontal cortex, central midline structures (ventromedial PFC and posterior cingulum) and parietal cortex during food anticipation. In addition, greater activation in the secondary gustatory cortex (posterior lOFC) but not in the primary gustatory cortex (insula/ frontal operculum) could be observed during the receipt of food reward. Finally, the BMI was positively correlated with bilateral lOFC activation during the receipt of food reward.

The VS is considered a central region of the reward system, responding to the anticipation of different types of reward (Knutson et al., 2001a; Rademacher et al., 2010; Sescousse et al., 2013). Particularly, the overall saliency of a stimulus is related to the activity in the VS (Zink et al., 2004). Correspondingly, we observed a linear increase in VS activation to increasing food reward. Furthermore, we observed bilateral anterior insula activation during the anticipation of food. The anterior insula is known to be related to interoceptive awareness including craving elicited by different drug cues (Craig, 2009). Since the gustatory cortex is located in the insula and overlying frontal operculum (Craig, 2009; Veldhuizen et al., 2011), this observation likely represents conscious urges for food rather than primary gustatory processing. Additionally, we found dorsal thalamic activation during the anticipation, but not the
receipt, of food reward which is in line with previous studies (Komura et al., 2011; Knutson and Greer, 2008), indicating that dorsal thalamus activation reflects increased arousal and attention primarily to reward predicting cues. In sum, the observed activations indicate a recruitment of the general incentive salience network during the anticipation of food related stimuli (Berridge and Robinson, 1998).

Receipt of food rewards recruited the bilateral lOFC and frontal operculum, indicating that the receipt of abstract food related stimuli recruits similar regions than those commonly observed during the processing of gustatory properties of food (Rolls, 2008; Small, 2012; Veldhuizen et al., 2011). Furthermore, receipt of food reward was associated with bilateral activations of the lPFC. This is in accord with a previous meta-analysis which found right lPFC activations during the processing of taste reward (Sescousse et al., 2013). The lPFC plays a central role in processes of attention, working memory, and inhibitory control (Owen et al., 1999; Tanji and Hoshi, 2008). In healthy subjects, suppression of cue induced food craving was associated with increased activation in the lPFC (Siep et al., 2012). Furthermore, repetitive transcranial magnetic stimulation of the left lPFC was found to reduce the desire for food consumption (Uher et al., 2005). Thus, bilateral lPFC activations during the receipt of food reward may reflect the recruitment of inhibitory control processes.

We did not observe any significant activity in the mOFC during the receipt of food rewards. The mOFC is considered relevant for the evaluation of food reward (Gearhardt et al., 2011). Although some previous studies found mOFC activation during the processing of different types of reward including food rewards (Peters and Büchel, 2010), we only observed activation in the lateral part of the OFC. Our findings are in line with previous studies, showing that the intensity of lOFC activation during food ingestion is closely related to subjective pleasantness ratings.
(Kringelbach et al., 2003). Hence, our results indicate that the neural coding of the receipt of abstract food rewards specifically involves the primary and secondary gustatory cortex, rather than common reward related brain regions (i.e. mOFC).

As expected, both tasks activated the VS during the anticipation of rewards. Heightened VS activity in response to different kinds of reward such as food, money, erotic stimuli or even pleasant music (Menon and Levitin, 2005; Sescousse et al., 2010; Simon et al., 2014; Simon et al., 2010) confirm the role of the VS as a site of modality-independent computation of reward-related salience (Metereau and Dreher, 2013). Differences in VS activation found in previous studies could be due to the fact that the delivery of monetary rewards often depends on the participants’ motor reactions, whereas food rewards are often delivered in a passive manner (Sescousse et al., 2013). Here, we compared for the first time food with monetary rewards in the same context. Our results are in line with a study by Sescousse and colleagues (2010), who compared erotic with monetary rewards using the same task and also failed to observe significant differences in VS activity between tasks.

Furthermore, a cluster of activation located in the posterior cingulate cortex (PCC) was stronger activated during the anticipation of food rewards compared to monetary rewards. Activation of the PCC was found to be increased in response to high caloric visual food and taste stimuli (Siep et al., 2009) and has been linked to craving for different types of reward. Together with the medial PFC, the PCC forms a system of cortical midline structures involved in self-related cognitions and attentional tracking of self-relevant stimuli (Northoff et al., 2006; Small et al., 2003). This is further corroborated by higher activation in the angular gyrus of the parietal cortex which is an additional region involved in self-awareness and attentional processing (Seghier, 2013). Therefore, compared to monetary reward, food reward seems to be
associated with increased self-referential processing (i.e. food is to a greater extent analyzed in relation with one’s self (Brewer et al., 2013)). Additionally, the observation of a stronger activation in the posterior IOFC during the receipt of food rewards is in line with previous research stating that innate rewards recruit phylogenetically older regions in the posterior OFC whereas learned reward associations recruit phylogenetically newer regions in the anterior OFC (Kringelbach and Rolls, 2004). Lateral portions of the OFC are strongly interconnected with the primary gustatory cortex, and both brain regions are involved in the conscious experience of pleasure from palatable foods (Friederich et al., 2013).

The positive relation between activations in the bilateral IOFC and BMI may demonstrate the impact of neural responsiveness to food stimuli on eating habits. This observation is in line with some, but not all, previous studies: Hyperactivation in the IOFC while viewing pictures of high-calorie food has been found to be associated with higher BMI levels (Rothemund et al., 2007; Stoeckel et al., 2008; Yokum et al., 2011). However, another study found an inverse relation between BMI and activation in the right IOFC during the observation of food pictures (Killgore and Yurgelun-Todd, 2005). Since activity in lateral orbitofrontal regions has been linked to the current reward value of tastes (Rolls et al., 2003; Simmons et al., 2005), and seems to be increased by hunger (Porubská et al., 2006), one may argue that a stronger activation in those regions contributes to increased food intake. Indeed, there is preliminary evidence that future increases in BMI are related to increased activity in the IOFC, but to a reduced sensitivity in the common reward circuitry, specifically the VS (Stice et al., 2010; Yokum et al., 2011).

A possible limitation of our study may arise from the use of a task employing abstract food stimuli that may differ from tasks that include the receipt of actual taste stimuli
during an fMRI scan. However, since the primary aim of the present study was to compare food- with monetary related reward stimuli, we used the same task design for both conditions. Furthermore, as all participants were women, results should be generalized to men with caution. Finally, all our participants had a BMI within the normal weight range. Therefore, it remains unclear whether our findings are also valid for overweight and obese people.

Taken together, the present findings support the notion that food and monetary reward processing engage both common and distinct brain networks. The direct comparison of both types of reward, assessed with the same task design in the same sample, revealed stronger activation in a brain network subserving self-referential processing. During the receipt of food reward, higher activation was observed in phylogenetically older portions of the IOFC. This underlines the posterior-anterior distinction of innate and learned reward processing in the OFC. Finally, the present task proved valid that both the anticipation and receipt of food reward can be reliably measured using abstract stimuli. Therefore, the present fMRI task offers a valuable method to further investigate brain circuits underlying reward processing in obesity and eating disorders.
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Figure 1: Graphical depiction of the money (MID task) and food incentive delay task (FID task). Cues representing possible reward outcomes (1 EUR, 20 cents and 0 EUR/10 snack points (SP), 2 SP and 0 SP, respectively) and task structure of the MID/ FID. Participants were presented with a cue stipulating the amount of money/ SP they could win if they reacted correctly during the following discrimination task. Immediately after target presentation, participants were informed about the amount of money/ SP they had won during the trial and their cumulative total win so far.

Figure 2: (a) Significant brain activation during the contrast *anticipation_high* for the FID task. The threshold was set at *p*<0.001 uncorrected, with a cluster-defining threshold of 5 voxels for illustrative purposes. The blue circles denote the localization of the VS ROIs. SPM t-map was rendered on a T1-weighted template image (coronal slice, *y* coordinate = 4 mm) supplied with mricon (Colin brain). (b) Bar chart depicting percent signal change from the right VS during different anticipation conditions for the FID task. (c) Bar chart depicting percent signal change from the left VS during different anticipation conditions for the FID task. Error bars depict SEM.

Figure 3: (a) Significant brain activation during the contrast *receipt_high* for the FID task. The threshold was set at *p*<0.001 uncorrected, with a cluster-defining threshold of 5 voxels for illustrative purposes. The blue circles denote the localization of the OFC ROIs. SPM t-map was rendered on a T1-weighted template image (transversal slice, *z*-coordinate = -8 mm) supplied with mricon (Colin brain). (b) Bar chart depicting percent signal change from the right IOFC during different snack points (SP) receipt conditions. (c) Bar chart depicting percent signal change from the left IOFC during different SP receipt conditions. Error bars depict SEM. (d) Correlation between percent signal change extracted at the right IOFC during the contrast *receipt_high*
and BMI. (e) Correlation between percent signal change extracted at the left IOFC during the contrast receipt_high and BMI.

**Figure 4:** (a) Whole brain analysis depicting areas with stronger activation during the FID then MID task during the contrast anticipation_high. (b) Whole brain analysis depicting areas with stronger activation during the FID then MID task during the contrast receipt_high. The threshold was set at p<0.001 uncorrected, SPM t-maps were overlaid on a T1-weighted template image (transversal slice, z-coordinate = -12mm) supplied with mricron (Colin brain).
25x12mm (600 x 600 DPI)
Table 1: Whole brain results for the anticipation/ receipt of primary and secondary rewards.

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<td>-6</td>
<td>***</td>
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<tr>
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<td>35</td>
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<td>***</td>
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<td>4.36</td>
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Table 1: All results p<.001 voxelwise uncorrected, except: * = voxelwise FWE corrected p<.05, ** = Cluster level p<.001 uncorrected, *** = Cluster level FWE corrected p<.05. BA; Brodmann area, L; left, R; right, MNI; Montreal Neurological Institute, ACC; anterior cingulate cortex, OFC; orbitofrontal cortex.
Table 2: Group maximum t-values and MNI-coordinates of all activation foci found during the comparison of the MID and FID task.

<table>
<thead>
<tr>
<th>Area</th>
<th>Anticipation of high vs. no reward</th>
<th>Receipt of high vs. no reward</th>
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<tr>
<td></td>
<td>t-value at peak voxel</td>
<td>MNI-coordinates</td>
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<td>x      y    z</td>
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</table>

Table 2: All results p<.001 voxelwise uncorrected, except: * = voxelwise FWE corrected p<.05, ** = Cluster level p<.001 uncorrected, *** = Cluster level FWE corrected p<.05. BA; Brodmann area, L; left, R; right, MNI; Montreal Neurological Institute.