Double Dissociation of Exteroceptive and Interoceptive Feedback Systems in the Orbital and Ventromedial Prefrontal Cortex of Humans

Elisabeth Hurliman, Jennifer C. Nagode, and José V. Pardo

The primate ventral prefrontal cortex contains two densely interconnected subregions: a lateral/orbital cortex processing primarily sensory/exterceptive information, and a ventromedial cortex processing primarily visceroappetitive/interoceptive information. These regions have major afferents and efferents to other associative cortices. The organization of these structures leads to an hypothesized role in feedback processing. We use neuroimaging to test this model, defined so far mostly through anatomical studies.

Healthy volunteers were trained operantly on a transitive inference task (A > B, B > C \(\Rightarrow\) A > C) requiring flexible manipulation of feedback to solve. Two groups of subjects learned an arbitrary face hierarchy, one adjacent face pair at a time; each group received either visual/exterceptive (“XXXX”) or visceroappetitive/interoceptive (fruit juice) feedback for correct responses to adjacent face pairs. After task acquisition, the subjects were tested on novel stimulus pairs (i.e., nonadjacent, TEST) derived from the acquired hierarchy. The TEST condition required transitive inference. No feedback was provided during TEST.

Brain activity during TEST in the group trained with visual/exterceptive feedback increased in the orbital prefrontal cortex and decreased in the ventromedial prefrontal cortex. In contrast, brain activity during TEST in the group trained with visceroappetitive/interoceptive feedback decreased in the orbital prefrontal cortex and increased in the ventromedial prefrontal cortex.

These results provide functional evidence, consistent with previous anatomical studies, for two major feedback systems in human ventral prefrontal cortex: a lateral system specialized for exteroceptive information and a medial system specialized for interoceptive information. Although highly interconnected, there is a double dissociation of function between these networks in healthy humans.

Key words: feedback processing; ventral prefrontal cortex; PET neuroimaging; transitive inference; associative learning; double dissociation

Introduction

The ventral prefrontal cortex (VPFC) participates in diverse cognitive functions, including representation of primary and secondary sensory information, associative learning, flexible updating of learned associations, and sensory-specific satiety (for review, see Rolls, 1996). It also seems critically involved in decision making and goal-directed behavior (Bechara et al., 1997; Elliott et al., 2000; Winstanley et al., 2004). A common theme across the cognitive functions of this region is their dependence on feedback.

Anatomical data indicate that the primate VPFC functions in feedback processing. Barbas and Rempel-Clower (1997) found that the VPFC projects to the more laminated dorsal prefrontal cortex in a feedback pattern. In turn, dorsolateral prefrontal regions project to ventral regions in a feedforward pattern. Öngür and Price (2000) and Öngür et al. (2003) proposed that two distinct, yet interacting, networks exist within VPFC: a lateral sensory network processing exteroceptive information [orbital prefrontal cortex (OPFC)] and a medial visceral network [ventromedial prefrontal cortex, (VMPFC)] processing interoceptive information such as visceral motor output.

Compared with the broad literature concerning exteroceptive/sensory systems, less is known about interoception. Interoception is the sense of the physiological condition of the body including, but not limited to, the viscera (for review, see Craig, 2002). Small-diameter afferents pass information of the physiological status of tissues on to higher cortex. Interoception subsumes multiple processes, such as nociception, thermoregulation, pH balance, blood pressure, thirst, swallowing, skeletal muscle stretch, neuroimmune interactions, etc. We previously noted large cortical activations when attempting to taste deionized, distilled water reflecting both intraoral somatosensory processing as well as fluid stimulation of the tongue and oropharynx (Zald and Pardo, 2000b). Interoceptive information inherently
has an affective component that provides motivation; likewise, water is a primary, unconditioned reinforcer. Although the administra-
tion of fluids has a chemoreceptive component attributable to taste, the value of such a stimulus directly depends on the physiological needs of the body and has relevance for maintenance of homeostasis and survival; these latter aspects group such stimuli within interoception (Craig, 2002).

In the neuroimaging literature, the term “feedback” is used loosely to refer to information after a response. However, in the present study, feedback is defined more narrowly as information provided after a response critical for responding correctly during incremental/operant learning. Feedback, whether exteroceptive or interoceptive, is processed through circuits that decode information about the feedback stimulus as well as assign relative value to the response.

Here, we probe the functional aspects of the anatomically hypothesized VPFC feedback systems in humans using 15O-water positron emission tomography (PET) and a feedback-dependent cognitive activation task. We expect to see OFPC activations in the exteroceptive feedback condition; we expect to see VMPC activations in the interoceptive feedback condition. In addition, we are interested in exploring the functional nature of interaction between the two subsystems. In a serial model, exteroceptive sensory input reaches the OFPC interacting with the interconnected VMPC to produce visceromotor output: both networks should activate. In a parallel model, under constraints of limited capacity for dual processing, reciprocal activations should occur (i.e., lateral activations would occur with medial deactivations and vice versa).

Materials and Methods

Experiment 1: visual feedback during training

Subjects. Ten subjects (mean age, 23 years; age range, 19–29 years; three female, seven male) without significant medical history were recruited from the local University of Minnesota campus. All were free of lifetime occurrence of major psychiatric disorders [per computerized Diagnostic Interview Schedule (DIS) screener] (Bucholz et al., 1991). Seven subjects were right-handed, two subjects were ambidextrous, and one subject was left-handed [Edinburgh Handedness Inventory (Oldfield, 1971)]. All subjects gave written informed consent as approved by the Institutional Review Committee and the Radioactive Drug Research Committee.

Cognitive activation task. For an initial description of this task, please refer to the work of Nagode and Pardo (2002). We designed a feedback-dependent cognitive activation task based on transitive inference (TI) (\(A \rightarrow B, B \rightarrow C \Rightarrow A \rightarrow C\)), with pictures of 10 faces arranged hierarchically along an arbitrary dimension (Nagode and Pardo, 2002). The underlying hierarchy was initially created at random but remained constant during each subject’s training, as well as across subjects. Two hierarchies of male faces and two hierarchies of female faces were created. Stimulus hierarchy was randomly assigned across subjects of a given gender. During training outside of the scanner, two adjacent faces of the hierarchy were displayed on a video screen (~50 cm from the subject) with the program MEL 2.0 (Psychology Software Tools, Pittsburgh, PA). The faces were ~7 × 13 cm black and white images presented in pairs. Subjects selected the face higher on the underlying hierarchy. Responses were made by pushing one of two keys on a response box within 3 s of face presentation (right response key pushed by right index finger indicates choice of right face as higher on hierarchy; left response key pushed by left index finger indicates choice of left face as higher on hierarchy). The first response for each face pair required a random guess, but future responses could rely on feedback for the face pair received previously. Trialwise feedback was provided in the visual modality (“XXXXX” appears under the correct face). Training continued until criterion (90% correct responses). The subjects learned the relationship (>) between the two faces in an adjacent pair incrementally based on feedback. For instance, one stimulus presentation during training could be face 2 presented on the left half of the monitor and face 3 presented on the right half. No nonadjacent faces in the hierarchy are presented during training (e.g., subjects during this portion of the task will never see face 2 together with face 4, 5, 6, etc.). At the end of training, once the entire hierarchy has been successfully acquired, this task is no longer feedback dependent.

During neuroimaging, subjects solve the adjacent pairs condition as a CONTROL condition (e.g., presentation of face 2 and 3, face 3 and 4, etc., from the underlying hierarchy of 10 faces) and a nonadjacent pairs condition (TEST) as experimental condition. During TEST, subjects solve the task by identifying in the absence of feedback the relationship between faces in novel pairs generated from nonadjacent faces from the hierarchy (TEST); e.g., presentation of face 2 on one half of the screen together with face 4 on the other half of the screen, 2 and 5, 2 and 6, etc., but never 2 and 3 or any other adjacent faces from the underlying hierarchy). In TEST, subjects were asked to select the correct one of two nonadjacent faces generated at random. The TEST pairs were allowed to differ in the degree of separation (i.e., symbolic distance) in the underlying hierarchy. First and last faces of the hierarchy were not used. No feedback was provided during TEST. Because TEST contains no feedback, the only way to respond correctly is through inference based on feedback received during CONTROL (adjacent faces).

Selection of the adjacent pairs CONTROL condition was based on several factors. First, this task offers the needed comparison condition for experimental activation tasks that tap the VPFC. This focus on comparison conditions is needed because the VPFC does not lend itself to an eyes-closed rest condition because of its high tonic resting activity. Second, this task mimics closely the cognitive demands from the TEST condition yet is much less dependent on using previously received feedback to solve it successfully. Although CONTROL provides a close match to the cognitive demands of the TEST condition, this match is not absolute. There is a small difference in demands to working memory defined more broadly than just keeping information on-line insofar as TEST requires an inference judgment, whereas CONTROL does not. However, a previous study from our laboratory, which directly manipulated visual working memory for faces, did not find activations of interest identified in the current set of experiments. In the working memory study, a direct comparison was made between memory for faces (high working memory demands) to a control task of scrambled faces (few if any working memory demands) (Kuskowski and Pardo, 1999). Similarly, Rushworth et al. (1997) did not find that primes with inferior convexity lesions (i.e., removal of lateral area 47/12) had any significant decrements on a visual working memory task.

Overall, the TI paradigm under study has several advantages. It is based on a widely used behavior paradigm, i.e., incremental, associative learning, which warrants additional study. It permits the use of several types of feedback (e.g., cognitive, appetitive). Performance of the control task is not stimulus bound, i.e., correct performance requires the use of previous feedback; thus, sensory confounds from actual feedback administration can be controlled. The paradigm contains a feedback-independent comparison condition (adjacent faces) once the task is well learned. This is of particular importance for studying VPFC function because this region has high tonic activity at rest. Thus, comparison with a resting state will obscure task-dependent VPFC activation. The paradigm provides a route around the use of resting brain states.

PET scanning. Images were acquired using an ECAT 953B scanner (Siemens, Knoxville, TN) in three-dimensional mode with septae retracted. A two-dimensional transmission scan was used for attenuation correction. H2-15O PET was used to measure whole-brain activation (31 slices, 3.5 mm slice separation) with uniform signal recovery. A catheter provided venous access for tracer injection. A vacuum-molded pillow was used for head stabilization. Slow-bolus (30 s) intravenous injections of H2-15O (9.25 MBq or 0.25 mCi/kg, initially) were administered at the beginning of each trial. Acquisition scans (90 s) for each of the following conditions were collected: resting with eyes closed; performing a sensorimotor control task (random button press while viewing unfamiliar face pairs); selecting the correct face from an adjacent face pair (CONTROL); and selecting the correct face from a nonadjacent face pair (TEST). No feedback was provided during any of these trials. Each condition was followed by a 10 min interscan interval to allow isotope decay.
Data analysis. In-house software was used to align the transmission scan and emission scans. Images were reconstructed with filtered back projection using a Hanning filter. Final image resolution was 9 mm full-width half-maximum. Estimated regional cerebral blood flow (rCBF) is based on normalized tissue radioactivity to 1000 counts. Automated software was used to assess motion artifacts, and scans were excluded if motion exceeded 2 mm or 2° during a scan session (Mionoshita et al., 1994). Nonlinear warping normalized the anatomy to a standard stereotactic atlas (Talairach and Tournoux, 1988). Scans from each condition were averaged across subjects for pixelwise group analysis. Significant activations were defined as those equal or exceeding a Z value of 3.1 (p < 0.001). In addition, individual subtraction images were obtained for region-of-interest analysis. Figures of functional images were produced with in-house software [iv (Lee and Pardo, 2000)] and BrainTools software [mri3dX; K. Singh, Aston University, Birmingham, UK; http://www.aston.ac.uk/lhs/staff/singhkd/mri3dX/mri3dX.jsp].

Experiment 2: interoceptive feedback during training
Subjects. Seventeen subjects were recruited initially from the local university community and gave written informed consent. Three subjects were excluded because of equipment malfunction. One subject was excluded because of severe motion artifacts. Two subjects were excluded because of failure to reach the learning criterion of 90% correct. The final sample consisted of 11 subjects (four female), with a mean age of 26 years (range, 19–45 years). This final sample was closely matched to our sample from experiment 1 on demographic variables, including age, sex, and race. All were free of lifetime prevalence of major psychiatric disorders (per computerized DIS screener (Bucholz et al., 1991)]. None had major medical illness. Ten subjects were right-handed, and one was left-handed. No subject had participated in experiment 1 with visual/exteroceptive feedback. To control for satiety effects, subjects who arrived at their training session more than 2 h after their last meal were provided a snack. No subject felt hungry or thirsty at the start of training.

Cognitive activation task. Training on T1 with visceralappetitive/interoceptive feedback occurred until performance reached 90% correct. Fruit juice was selected for feedback based on subject palatability. An automated pump system controlled by MEL 2.0 software (Psychology Software Tools) delivered 0.25 ml of fruit juice via a plastic tube into the subject’s mouth for a correct response. Fruit juice was selected as the interoceptive feedback stimulus because of its extensive oropharyngeal processing beyond sensorimotor information (salivation, temperature, and viscosity), its ease of implementation in operant training, and its strong motivational component. To ensure that the juice was perceived as palatable, subjects reported any decrease in palatability during rest breaks every 10 min. The juice was changed according to the subject’s preference. We note that this interoceptive feedback condition differs from the exteroceptive feedback condition in terms of reward properties. However, interoceptive feedback processing has an inherent hedonic component that rules out any attempt to stimulate interoceptive feedback processing independently of rewarding (either pleasant or aversive) properties of the stimulus (for review, see Craig, 2002). The reward properties were minimized by using satiated subjects. Also, the repeated administration of the same fruit juice likely led to habituation in primary taste cortex (no anterior insula activation occurred during training with fruit juice), as well as sensory-specific satiety.

The CONTROL condition was matched in every way to the TEST condition, except the stimulus pairs were adjacent and had been acquired previously. Of note, CONTROL, like TEST, had no feedback. In addition to the advantages of the CONTROL condition reported for experiment 1, this comparison (TEST − CONTROL) avoids the confounding effects of visceral sensory processing in frontal secondary association cortices. As mentioned for experiment 1, one potential cognitive difference beyond feedback processing is the slightly increased demands on working memory in the TEST condition because of the required inference judgments. However, in this set of experiments, we included a sensorimotor control condition. The comparison (TEST − SENSORIMOTOR CONTROL) did not show significantly more active medial VPFC in the TEST condition, which has significantly greater working memory demands than the SENSORIMOTOR CONTROL of passively viewing faces while making random button pushes.

PET scanning and data analysis paralleled the procedures described for experiment 1.

Results
Behavioral results
Mean correct performance in experiment 1 for TEST was 98% and for CONTROL was 96%. Mean correct performance in experiment 2 for TEST was 95% and for CONTROL was 93%. Percentage correct responses were transformed to arcsine values to achieve a normal distribution. Independent sample t tests showed significant differences between TEST − CONTROL across feedback groups (t19 = 2.37; p = 0.029). The difference may reflect technical differences in training necessitated by the different feedback conditions. Nevertheless, both feedback groups performed above our performance criterion of 90% correct. More importantly, performance during TEST was not significantly different across feedback groups (t18 = 0.462; p = 0.650).

Whole-brain pixelwise subtractions: reciprocal lateral and medial VPFC activations
Figure 1 demonstrates the reciprocal changes that occur in OPFC and VMPC during the two feedback conditions. OPFC regions significantly activate in the contrast TEST − CONTROL for the visual/exteroceptive feedback condition (Fig. 1, Table 1). These networks arise bilaterally. In contrast, a trend toward OPFC deactivation emerged for the visceroappetitive/interoceptive feedback condition (coordinates, 39 mm, 35 mm, −9 mm; Z score of −2.6). The VMPC deactivates for the visual/exteroceptive feedback condition (Table 2) and activates for the visceroappetitive/interoceptive condition (Fig. 1). Of note, one of the highly deactivated VPFC regions in the visual/exteroceptive feedback condition coincides closely in location with a region highly activated in the visceroappetitive/interoceptive feedback condition. Thus, medial and lateral VPFC activity found in whole-brain group analyses followed a reciprocal pattern against external and internal feedback modalities.

In contrast, in the visceroappetitive/interoceptive feedback condition, VMPC was activated, not OPFC (Table 3, Fig. 2). Additional significant medial activations were found for the visceroappetitive/interoceptive feedback condition in the parahippocampal–amygdaloid transitional area and ventral temporal area/substantia nigra (VTA/SN) (Table 3, Fig. 2). Note that the anatomical labels and Brodmann areas (BA) should be considered approximate because of inherent limitations in resolution associated with PET imaging.

Although the observed changes in activity in the visceroappetitive/interoceptive feedback condition generally were lower than those in the visual/exteroceptive feedback condition, activity changes in the VPFC followed our initial hypotheses in location and pattern of activation. In addition, despite the slightly lower magnitude of activity changes in the visceroappetitive/interoceptive feedback condition, activations and deactivations were highly significant when considering resolution elements in our predefined VPFC region of interest.

Region-of-interest analysis: lateral OPFC, medial VPFC, and amygdala
Additional evidence for the double-dissociated interaction between the two ventral prefrontal systems emerges in a repeated measures ANOVA between the two feedback types (visual/ex-
terceptive and visceroappetitive/interoceptive) and three regions of interest (VMPFC, OPFC, and amygdala) (Fig. 3). Peak pixel activations based on the actual data set were used to delimitate regions of interest. We selected the amygdala as an additional region of interest because of its well established, dense neural connections with ventral prefrontal areas (Porrino et al., 1981). The VMPFC and amygdala change activity together. Mean activity in the exteroceptive feedback condition shows a decrease in both amygdala and VMPFC regions and an increase in the OPFC region. The opposite pattern emerges in the interoceptive feedback condition, because the amygdala and VMPFC regions show increased rCBF and the OPFC exhibits decreased rCBF (Fig. 3, Table 4).

In addition to the data-driven region-of-interest analysis, a global brain analysis was conducted in SPM99 (Wellcome Department of Cognitive Neurology, London, UK) for interactions between feedback conditions and the ventral prefrontal regions of interest. Results indicate significantly medial VPFC activation in the interoceptive feedback condition (Z score of 2.5), and significant OPFC activation was found in the exteroceptive feedback condition (Z score of 2.7) (Figs. 4, 5).

Discussion

Overview of results

Results of both experiments suggest the VPFC contains two feedback-processing systems, consistent with hypotheses derived from anatomical studies (Barbas and Rempel-Clower, 1997; Öngür et al., 2003). One subsystem, situated laterally in the VPFC, preferentially processes information from the external environment; the other subsystem, situated medially, preferentially processes information from the internal/bodily environment.

In addition, our data reveal that the two feedback systems function in a double-dissociated manner. Inference based on review of visual/exteroceptive feedback history activates OPFC and deactivates VMPFC. Conversely, inference based on review of visceroappetitive/interoceptive feedback history activates VMPFC and deactivates OPFC. The two VPFC systems thus seem poised toward competition.

Considerably less afferent sensory input exists to VMPFC than to OPFC. Taste feedback could activate OPFC regions such as posterolateral orbital cortex, and VMPFC regions may primarily process monitor visceromotor output and participate in decision making in this manner (Öngür and Price, 2000). Here, a functional double dissociation between VMPFC and OPFC would necessitate an inhibitory feedback loop. Visceral information travels via spinal, facial, glossoparyngeal, and vagal afferents before reaching the brainstem and cortex (Ceppetto and Chen, 1990). Fine afferents carrying interoceptive information reach hypothalamus and periaqueductal gray (Craig, 2002), which in turn are connected to VMPFC and limbic regions. Via reciprocal connections from amygdala to VMPFC (Price, 2003), interoceptive information may reach VMPFC without traveling through OPFC first. Although the number of afferents for exteroception far outnumber those for interoception, visceral information is not passed on in an undifferentiated manner (Jänig, 1996; Vaitl, 1996).

The current study presents novel findings regarding two double-dissociated VPFC feedback systems. Previous studies examined different functions of VPFC subregions with one feedback type. O’Doherty et al. (2000) studied monetary feedback. Although money is an external stimulus, it is a potent reinforcer
because it becomes closely associated with food/shelter, comfort, and pleasure. The feedback systems operating on monetary rewards and punishments require additional dissection.

Experimental issues
The selection of the comparison TEST — CONTROL was motivated by several factors. First, high tonic resting activity of the VPFC required a nonresting comparison task (Gusnard and Raichle, 2001; Mazoyer et al., 2001). Second, studying feedback processing without confounded sensory aspects requires a feedback stimulus-independent task during scanning. During TESTING, no feedback is administered, yet only with review of previously received feedback during scanning can the TESTING condition be solved. CONTROL during PET scanning (i.e., after a long training phase) is overlearned and feedback independent.

Working memory demands differ across TESTING and CONTROL, but this does not explain the current activations. When examining another comparison (TEST — SENSORIMOTOR CONTROL, i.e., few working memory demands during passive viewing of faces and random button pushes) for the interoceptive feedback group, no VMPFC activations emerge. This result is consistent with Rushworth et al. (1997) and Kuskowski and Pardo (1999), who have used a direct visual working memory task using face stimuli.

We implemented several experimental controls to keep constant rewarding properties of fruit juice feedback. First, we attempted to ensure that volunteers were satiated before task acquisition. Second, we adjusted the fruit juice per individual preference throughout training as needed. Third, we did not provide interoceptive feedback during imaging to control for possible habituation effects to feedback for well known face pairs. Nevertheless, the overall smaller activations observed in the interoceptive feedback condition may be partially explained by some satiety effects and changes in reward value across the duration of training.

Cognitive implications
The exact nature of the cognitive operations within VPFC remains undetermined. This region underlies encoding and flexible updating of stimulus–response associations (Rolls et al., 1994; Passingham et al., 2000; Nagahama et al., 2001). The temporal resolution of PET prevents more detailed study of time-dependent VPFC activity. Our preliminary functional magnetic resonance imaging (MRI) block design studies suggest that a buildup of blood oxygenation level-dependent signal occurs in OPFC during training.

This study does not clarify the nature of the representation of feedback-dependent associations between stimulus and correct response. Although not the focus of this study, the same precuneus region activated during TESTING across both studies. Thus, visual codes may be used in the hierarchy representation. Likewise, little is known about the representation of the relational operator (greater than, >) and its relationship to affective/motivational processes. Activity in the amygdaloid region, situated medially in periamygdaloid cortex/basal forebrain regions, varied along with the VMPFC across feedback types. Such activation is consistent with projections from medial aspects of the amygdala to hypothalamus, and midbrain, participating in the regulation of visceral functions (for review, see Price, 2003). Of note, subjects were neither thirsty nor hungry at the time of training or TESTING. Therefore, the amygdaloid activity probably is not coding the valence of the feedback but more likely the integration of autonomic interoceptive processes related to the feedback.

The VTA/SN also activated during TESTING after training with visceroregulatory feedback. Connections between VMPFC and VTA/SN are well established (Porrino et al., 1981; Öngür and Price, 2000; Tzschentke and Schmidt, 2000), and its role in feeding as part of the mesolimbic dopamine pathway has been identified.

### Table 3. Significant ventromedial activations (VMPFC) in TEST — CONTROL, interoceptive feedback during training

<table>
<thead>
<tr>
<th>Region</th>
<th>Coordinates</th>
<th>Z score</th>
</tr>
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<tbody>
<tr>
<td>Extended amygdala</td>
<td>34</td>
<td>-6 -11</td>
</tr>
<tr>
<td>Ventral segmental area/substantia nigra</td>
<td>10</td>
<td>-19 -9</td>
</tr>
<tr>
<td>Gyrus rectus (GR)</td>
<td>14r</td>
<td>6 21 -16</td>
</tr>
</tbody>
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The region in parentheses is per the three-dimensional probabilistic atlas (Chiavaras et al., 2001). Brodmann areas (BA) are assigned per Price (Öngür et al., 2003), mapped onto the atlas of Talairach and Tournoux (1988).
(Korotkova et al., 2003; MacDonald et al., 2003). Recent neuroimaging studies are beginning to show that both VMPFC and VTA/SN are important in human food consumption and obesity (J. V. Pardo, S. A. Sheikh, B. R. Ritberg, and D. E. Adson, unpublished observation), consistent with a role in viscerovipettive/interoceptive processing. Stimulation of OPFC has multiple autonomic effects, suggesting that the VPFC is a major cortical representation of the vagus nerve (for review, see Fuster, 1989). Unfortunately, the resolution prevents more definitive identification of such structures activated during TEST.

How can fruit juice, a sensory stimulus, activate medial/interoceptive systems without activating lateral/exteroceptive systems? Typically, activations in primary and secondary association cortices for flavors, tastes, and smells are not observed with repeated stimulation (Small et al., 1999; Zald and Pardo, 2000a). As expected, we do not find primary or secondary taste cortex activations with the fruit juice stimulus delivered over >1 h. Thus, the observed activations do not represent the sensory/discriminatory stimulus component but its intrinsic value.

Could a difference in rewarding properties between the two feedback types be the main reason for the different activations? A follow-up study that directly manipulated the “wanting” component of reward clarifies this. There, fasting subjects acquired the TI task with interoceptive feedback. All subjects were fed before PET scanning. In the follow-up, more anterior VMPFC activations emerged (fasting, y = 32 mm; satiated, y = 21 mm), consistent with other studies that observed anterior VMPFC activation with rewarding stimuli (Gottfried et al., 2003; Kringelbach and Rolls, 2004). Furthermore, although we observe activations in VTA/SN, there is no clear consensus in the literature regarding the structures underlying aspects of reward processing (“wanting” vs “liking” (Wyvell and Berridge, 2000)). In the follow-up experiment, we observe additional strong activations in anatomical regions typically associated with reward, such as ventral striatum (coordinates, −21 mm, 14 mm, −9 mm; t = 4.5 in a within-group comparison). It is possible that VTA/SN activations observed in the current study represent “liking” or the hedonic component of interoceptive stimuli. VPFC activations in the current studies emerged only in a limited portion of this extensive frontal region. Functional specificity likely exists within VPFC. In addition to sensory association cortices for taste in the insula and anterior cingulate, the VPFC is known to participate in sensory taste processing (Kobayashi et al., 2002; Rolls et al., 2003; de Araujo et al., 2004). It also participates in the flexible updating of associations to adjust to new contingencies (Rolls et al., 1994; Freedman et al., 1998; Baxter et al., 2000). Furthermore, some VPFC region may be crucial in predicting reward value (Rolls et al., 1989; Tremblay and Schultz, 1999; O’Doherty et al., 2000), and neurons seem to discriminate between reward values (Rolls et al., 1996; Passingham et al., 2000; Gottfried et al., 2003).

The functional double dissociation within VPFC feedback subregions suggests that exteroceptive and interoceptive feedback cannot be processed equally at the same time. Inappropriate activation of the medial subsystem may lead to deactivation of the lateral subsystem at the expense of exteroceptive feedback processing. Limited capacity for dual processing could have profound implications for cognitive disorders with impaired deci-
sion making in general and addiction in particular. Overemphasis of interoceptive stimuli could lead to vascrally biased decisions that neglect external/cognitive signals. Such bias may allow addicts to neglect long-term negative social and occupational consequences associated with drug use in favor of immediate visceral rewards or relief of negative visceral stimuli. Paulus et al. (2002) showed that methamphetamine-dependent subjects failed to activate this region (BA 10/11) during a two-choice task. Bechara and Damasio (2002) found differences in a VMPFC-dependent gambling task in addicted subjects. Preliminary data of several long-term cocaine users showed that these volunteers inappropriately recruited the VMPFC and deactivated the OPFC when solving our cognitive activation task under the visual/exteroceptive feedback condition (our unpublished observation). Implications thus exist for psychopathological conditions with impairment in decision-making attributable to overemphasis of interoceptive stimuli (Porrino and Lyons, 2000; Goldstein and Volkow, 2002), because adaptive functioning relies partly on appropriate recruitment of VPFC feedback regions.

References


Nagahama Y, Okada T, Katsumi Y, Hayashi T, Yamauchi H, Oyanagi C,


