

Anticipatory Cues Differentially Provoke *In Vivo* Peptidergic and Monoaminergic Release at the Medial Prefrontal Cortex

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Like primary reinforcers, the anticipation of reward ought to affect neurochemical release in brain regions, such as the medial prefrontal cortex (mPFC), which are associated with appraisal processes. To assess the neurochemical changes associated with anticipation, rats were exposed to the pairing of auditory (60-dB white noise), visual, and olfactory cues with the daily presentation of a palatable snack (Cue Relevant group). Rats of a second group were similarly trained, but for a 2-week period, the snack was no longer provided following cue presentation (Extinction group). In the third condition, the presentation of the snack and cues was uncorrelated (Cue Irrelevant group). Analyses of dialysates collected *in vivo* from the mPFC revealed that release of corticotropin-releasing hormone (CRH), gastrin-releasing peptide (GRP), and the 5-HT catabolite, 5-hydroxyindole acetic acid (5-HIAA), had increased bilaterally in response to the anticipatory cues, whereas DA release increased only within the right mPFC. In the case of CRH and GRP, these increases were also apparent in the extinction condition, despite the fact that behavioral arousal to the anticipatory cues (increased exploration, rearing, grooming, and vigilance) was only evident in the Cue Relevant condition. In contrast, the elevated DA and 5-HIAA were apparent exclusively in the Cue Relevant condition. Thus, CRH and GRP systems may serve to allocate salience and/or incentive reward value to biologically significant stimuli or reflect the emotional response to the anticipatory stimulus. The activity of DA and 5-HT neurons, in contrast, is more closely aligned with the cognitive appraisal of predictor stimuli.

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INTRODUCTION

Appetitive and aversive events provoke neurochemical changes within the central and peripheral nervous systems, and it might be expected that the anticipation of such events might also promote adaptive neurochemical responses. The medial prefrontal cortex (mPFC) may be particularly important in this regard, given its role in higher cognitive functions (Kalivas and Nakamura, 1999; Van Eden and Buijs, 2000). Indeed, increased mPFC dopaminergic (DA) and serotonergic (5-HT) activity are associated with both aversive (Bray, 2000; Horger and Roth, 1996; Sullivan and Gratton, 1999; Goldstein *et al.*, 1996) and appetitive processes (Bassareo and Di Chiara, 1997; Blackburn *et al.*, 1989; Feenstra and Botterbom, 1996; Jodo *et al.*, 2000;

Nakahara *et al.*, 2000; Pratt and Mizumori, 2001; Richardson and Gratton, 1996; Wise, 2000; Richardson and Gratton, 1998); Interestingly, accumbal DA utilization was more closely aligned with preparatory or anticipatory food-related stimuli than with ingestion itself (Blackburn *et al.*, 1989), although DA variations were induced by deviations of expected reward outcomes (eg delaying reward presentation or reducing reward duration) (Richardson and Gratton, 1998). Such effects are not unique to mesolimbic DA circuits, as hypothalamic 5-HT release was also evoked by both food-related cues and food consumption (Hoebel *et al.*, 1989, 1992; Schwartz *et al.*, 1990). Thus, 5-HT neurons may be important in an anticipatory capacity, possibly reflecting incentive motivation, emotional aspects of anticipation, or cognitive information processing.

Corticotropin-releasing hormone (CRH), which has typically been considered in the context of stress, fear, and anxiety (LeDoux, 2000; Davis, 1998), has been implicated in appetitive (ingestive) responses as well (Merali *et al.*, 1998). It would be of considerable interest to determine whether this peptide is also involved in anticipatory processes. Furthermore, gastrin-releasing peptide (GRP), which is activated with food ingestion and

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stimulates hypothalamic-pituitary-adrenal activity (Gibbs *et al*, 1979; Gibbs and Smith, 1988; Kent *et al*, 1998; Merali *et al*, 1999, 2002), is released at the central amygdala in response to both appetitive and aversive stimuli (Merali *et al*, 1998). Although CRH and GRP systems may functionally be interlinked (Merali *et al*, 2002), and receptors for both peptides are expressed within the mPFC (Kroog *et al*, 1995; Radulovic *et al*, 1998), limited information is available as to whether these peptides are associated with anticipatory processes.

The current investigation assessed *in vivo* release of DA, 5-HT, CRH, and GRP at the mPFC in relation to the 'anticipation' of a palatable snack. Considering that stressor-elicited DA activation at the mPFC is laterally biased (asymmetrical) (Sullivan and Gratton, 1998; Nielsen *et al*, 1999; Brake *et al*, 2000; Carlson *et al*, 1993), we assessed whether anticipatory monoaminergic and/or peptidergic variations were likewise laterally asymmetrical. Importantly, rather than assessing the effects of anticipatory cues in food-deprived animals, we did so in sated rats trained to anticipate a 'palatable snack', thereby circumventing the potentially confounding effects of food deprivation. Finally, in order to ascertain the specificity of the anticipatory neurochemical variations, neurochemical changes were also determined in rats initially trained to anticipate the snack, but then exposed to an extinction regimen.

MATERIALS AND METHODS

Subjects and Apparatus

A total of 45 male Long-Evans rats, weighing 275–300 g, were obtained from Charles River Inc (St Constant, Quebec). After arrival in the vivarium, and a mandatory quarantine period, animals were individually housed in custom-designed 'anticipation' cages (clear Plexiglas boxes, 35 cm × 25 cm × 34 cm with a wire mesh floor and a 'safe' cubicle of black Plexiglas 15 cm × 15 cm × 19 cm located in one corner of the cage). Cages were equipped with a speaker that delivered 60-dB white noise. Animals were fed *ad libitum* (except during training and testing) and housed in a quiet room, separated from other animals in the facility, in a controlled environment maintained at 21°C with 60% relative humidity. All experimental procedures followed the Canadian Council on Animal Care guidelines and were approved by the Research Ethics Committee of the University of Ottawa.

Training

Animals were randomly assigned to three groups and were exposed to their respective training procedures both before stereotaxic surgery and for 2 weeks following recovery. Briefly, animals were weighed each morning, during which their cages were cleaned and their food was removed, so that the presence of the experimenter did not become a conditioned stimulus. Over the next 2–3 h, the experimenter randomly entered and exited the room, mimicking the dialysate sample collection procedure that would subsequently be introduced. Rats in one condition (Cue Relevant group) were presented with compound 'anticipatory'

cues that comprised a 60-dB white noise emitted through a speaker located in the cage top coupled with the sight/odor of the snack (two 1.8 g squares of Graham Wafers) (Christie's Honeymaid[®]) placed at the top of each cage, but inaccessible to the rat. On each day (between 1100 and 1300), this procedure was repeated for variable time periods (5–40 min) prior to snack delivery, after which rats were given access to the palatable snack and the auditory cue was immediately terminated. These rats received the treatments during the 2-week period preceding surgery and for 2 weeks beginning 5 days after surgery. Rats in a second condition (Extinction group) were similarly trained (exposed to the cues and given access to the snack); however, after 4 weeks of initial training and the surgical procedure, the snack delivery was discontinued for the 2-week period prior to testing, although the anticipatory cues continued to be presented. Finally, rats in the third condition (Cue Irrelevant group) were presented with the compound stimulus and the snack, but the two were independent (uncorrelated) of one another. This procedure was undertaken for 2 weeks before and 2 weeks after surgery, with the latter commencing after the 5-day surgical recovery period. It will be noted that in well-trained animals, on presentation of a palatable snack rats immediately approached and consumed this food. Moreover, we have observed that when a palatable snack is presented to animals in their living cage, it did not elicit an anxiety-like reaction (although anxiety was provoked when the snack was offered to animals in unfamiliar surroundings) (Merali *et al*, 2003), and hence the neurochemical effects of this treatment in the present investigation could not be attributed to anxiety effects.

Surgery

Following the initial 2-week training period, rats were anesthetized using pentobarbital (60 mg/kg, intraperitoneal) with halothane supplement (1.5 l) as required, and placed in a stereotaxic instrument with level skull. Blunted ear bars were used in order to protect the rats' eardrums. Two 20-gauge guide cannulae, each containing a removable 24-gauge stainless-steel obturator, were implanted at the right and left mPFC. The placement coordinates (Paxinos and Watson, 1986) with level skull were antero-posterior = +3.2, lateral = ±0.8, and dorsoventral = -3.5. The guide cannula (which protruded from a custom-designed Delrin[™] pedestal) was then secured to the skull with four stainless-steel screws and acrylic dental cement.

In Vivo Microdialysis and Behavioral Testing

At 1 day prior to testing, animals were briefly anesthetized using halothane and the obturators (within the guide cannulae) were replaced with microdialysis probes. The concentric microdialysis probes (with 2 mm of active membrane; 250 µm outer diameter) of regenerated cellulose (6000 Da molecular weight cutoff; Spectrum Medical Industries) were positioned above the target site such that they rested within the guide cannulae. Each probe was secured with a retaining screw and connected via polyethylene tubing (Intermedic, Clay Adams, NJ) to a liquid swivel and a 2.5 ml infusion syringe (Hamilton) attached to

a pump (model 22, Harvard). Microdialysis probes were perfused overnight at a rate of 0.5 $\mu\text{l}/\text{min}$ with filtered KRB solution consisting of (in mM): K^+ 2.7, Na^+ 145, Ca^{2+} 1.35, Mg^{2+} 1.0, Cl^- 150, (pH 7.4) and BSA (0.1%) (Moghaddam and Bunney, 1989). On the following morning, animals were gently restrained and probes were lowered into the target position (within the mPFC), and the perfusion rate of the KRB was increased to 3 $\mu\text{l}/\text{min}$. Sample collection was initiated 1.5–2 h later. On collection, each sample (50–60 μl) was immediately frozen on dry ice and stored at -80°C until analysis by radioimmunoassay (RIA) and high-performance liquid chromatography (HPLC).

On the test day, the probes were continually perfused with KRB, and dialysates were pooled every 20 min throughout the experiment. After collecting five baseline samples, all rats were presented with the ‘anticipatory’ cues, as described earlier, for a 40 min period. Dialysate samples continued to be collected every 20 min. At the end of the ‘anticipatory’ period, rats of the Cue Relevant group were presented with and allowed to consume the snack, and two dialysate samples were collected during this 40 min period. For the two remaining groups, the snacks were removed from the top of the cage and discarded, but samples continued to be collected.

Radioimmunoassays

The detection and quantification of CRH was achieved through a solid-phase high-sensitivity adaptation/modification (Maidment and Evans, 1991) of the double antibody liquid-phase RIA originally described by Vale *et al* (1983). BN-LP(s) were detected using a similar solid-phase RIA (Plamondon and Merali, 1997). Briefly, protein A/G (Calbiochem-Novabiochem Corp., La Jolla, CA)-coated Immulon[®]-4 wells (Dynatech Laboratories, Inc., Chantilly, VA) were incubated with anti-CRH serum (rC70 kindly provided by W Vale) or anti-BN serum (α -BN2 kindly provided by Dr T Moody) for 2 h at 20°C . Samples, standards (reconstituted in the KRB solution with BSA (0.1%), ranging from 0.05 to 250 fmol/well or blanks, were incubated for 24 h at 4°C . Next, 25 μl assay buffer containing 5000–6000 cpm [^{125}I -Tyr⁰]rCRF (Amersham Canada Ltd, Oakville, Ontario) or [^{125}I -Tyr⁴]BN (iodinated in house, as per Salacinski *et al*, 1981) was added to each well and incubated for an additional 24-h period at 4°C . Finally, the wells were rinsed, separated, and their residual radioactivity was counted in a gamma-counter (Cobra[®] II Auto-gamma, Meriden, CT). A four-parameter logistic curve fit model was used for interpolation of the standard curves. The sensitivity of the assay was typically about 0.1 and 2 fmol/well for CRH and BN, respectively.

The specific anti-CRF serum used in the study recognized CRH_{1–41} and displayed negligible crossreactivity with other related peptides (Vale *et al*, 1983), including urotensin 1 and urocortin (data not shown). The BN antibody used in the RIAs recognized the C-terminal fragment of BN and strongly crossreacts with amphibian BN (100%) and certain mammalian BN-LPs, including GRP_{1–27} (110%) and GRP_{18–27} (neuromedin C or NMC (82%)), but only weakly with GRP_{1–16}, NMB-10, NMB-32, or substance P ($\leq 0.1\%$). We have shown previously that the major source of BN-LP

immunoreactivity from the hypothalamus is attributable to GRP (Merali and Kateb, 1993).

High-Performance Liquid Chromatography

Levels of DA and 5-hydroxyindole acetic acid (5-HIAA) (5-HT was below the detection limit) were determined using an HPLC system (Agilent Technologies, Waldbronn, Germany). Samples were injected (Agilent 1100 series Autosampler, Waldbronn, Germany) into the HPLC system equipped with a single-cell electrochemical detector (Antec Leyden Model Intro, Montreal, PQ, Canada) with an applied potential of 0.650 nA, filter of 1 s, and range of 0.1 nA/V. The separation of these analytes was achieved by their passage through an ESA, $4.6 \times 150 \text{ mm}^2$, 5 μm analytical column (Zorbax Eclipse XDB-C8, Agilent Technologies, Waldbronn, Germany). The mobile phase, consisting of 90 mM sodium dihydrogen phosphate (monobasic), 1.7 mM 1-octane sulfonic acid (sodium salt), 50 nM EDTA, 10% (200 ml/2l) acetonitrile, 50 mM citric acid (monohydrate), and 5 mM KCl (final pH = 2.4), was delivered at a flow rate of 1.0 ml/min. Quantification of the various analytes was accomplished by comparing their area under the curve to those of known external standards (calibrated at 2.5 pg/50 μl , 5.0 pg/50 μl , and 50.0 pg/50 μl) using a computerized Agilent ChemStation chromatography data acquisition system (Agilent Technologies, Mississauga, Ontario).

Behavioral Monitoring

During testing, animals were videotaped and their behaviors were analyzed over a 120 min time period (during baseline, ‘anticipation’, and ‘wafer’ sampling periods). The frequency of sleeping/inactivity, exploring (moving about or actively examining an area of the test environment), rearing, vigilance (alert watchful), and grooming were scored every 10 s. This time sampling procedure has been found, in our laboratory, to provide better than 90% agreement between independent raters.

Histology

At the end of the experiment, animals were deeply anesthetized using halothane and decapitated. Their brains were sectioned and stained for histological examination. Only animals with verified placement in the right and left mPFC were used in this study. Of the original 45 animal contingent, 23 animals had both cannulae correctly positioned within the mPFC, 11 had one cannulae correctly positioned (seven in the right hemisphere and four in the left), while 11 animals did not have either cannulae correctly situated.

Statistical Analysis

The microdialysis data were analyzed by averaging the first three baseline samples (denoted as 100%) and subsequent samples expressed as a percent of this baseline. This included the last two of the five baseline samples, the two samples taken during the anticipatory period, and the ensuing two samples when rats in the Cue Relevant group were permitted to consume the snack. The behavioral

responses (exploring, rearing, grooming, vigilance, and resting/sleeping) and the interstitial levels of CRH, GRP, DA, and 5-HIAA in the mPFC were analyzed by Huynh-Feldt corrected mixed measures ANOVAs with Treatment (Cue Relevant, Extinction, Cue Irrelevant) as the between-group factor and both Periods (Baseline, anticipatory period and consumption period) and Time (dialysates taken over the 20 min samples within each period) as the within-subjects factor. As cannulae placements were not uniformly accurate in both hemispheres, data from rats with only a single accurate placement were included in the analyses, and hence hemisphere (left vs right cannulae placement) was considered a between-group variable. In some instances, procedural difficulties precluded samples being collected and/or analyzed, and hence the degrees of freedom varied across the different amines and peptides. Where a significant interaction was present, follow-up comparisons of the simple effects were conducted by Newman-Keuls multiple comparisons ($\alpha = 0.05$).

RESULTS

Effect of the Anticipatory Cues and Subsequent Consumption (or not) of a Palatable Snack on the Animals' Behavior

Figure 1 depicts the frequencies of the animals' behaviors (Cue Relevant, $n = 11$; Extinction, $n = 10$; Cue Irrelevant, $n = 11$) in response to the presentation of the 'anticipatory' cues followed by the presentation (or not) of a palatable snack. The behavior of rats in the Cue Relevant group was characterized by alertness during the anticipatory period, whereas in the Extinction and Cue Irrelevant groups this behavioral profile was not evident, thus attesting to the

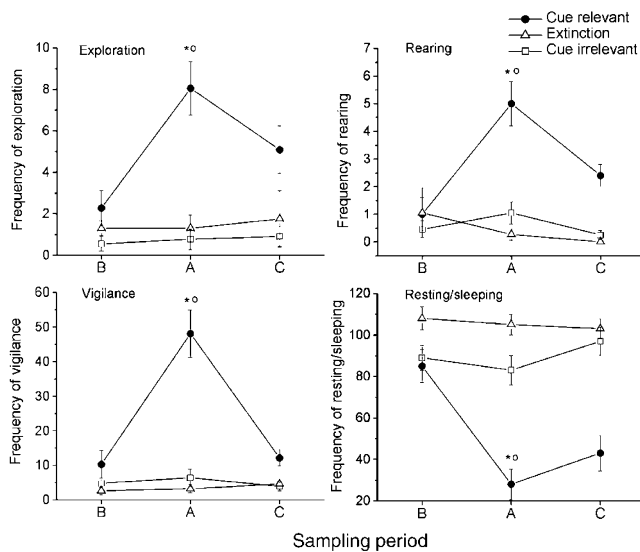


Figure 1 Frequency (mean \pm SEM) of exploration, rearing, vigilance, and resting/sleeping, using a 10 s time sampling procedure, among rats in the Cue Relevant, Extinction, and Cue Irrelevant conditions. Behaviors are presented during 20 min periods reflecting a baseline period (B), an anticipation period (A), and following snack presentation in the Cue Relevant group (C). * $p < 0.05$ relative to baseline (B5); $^{\circ}p < 0.05$ relative to Cue Irrelevant and Extinction groups.

apparent efficacy of the training procedure. Figure 1 shows the four most common behaviors (exploration, rearing, vigilance, and resting/sleeping) for each of the groups during the six sampling periods.

The frequency of resting/sleeping varied as a function of the Treatment Condition \times Period \times Sample interaction, $F_{4,58} = 13.33$, $p < 0.001$. During the baseline period, each of the groups spent the greatest proportion of time in a resting/sleeping state. In both the Extinction and Cue Irrelevant conditions, this continued through the remaining phases. In contrast, in the Cue Relevant group, the frequency of resting/sleeping declined markedly during the anticipation period, reaching statistical significance during the second portion of this period and the first portion of the consumption phase. During these times, rats in the Cue Relevant group rested/slept significantly less than did rats in the other two groups (see Figure 1).

The decline of resting/sleeping in the Cue Relevant condition was accompanied by an increase of active behaviors, including exploration, rearing, and vigilance. The ANOVAs revealed significant Treatment Condition \times Period \times Sample time interactions for these behaviors, $F_{4,58}^2 = 6.54$, 3.16, 3.84, p 's < 0.001 , respectively. The follow-up tests confirmed that during the anticipatory period, rats in the Cue Relevant group exhibited more exploration, rearing, and vigilant behavior relative to baseline. Moreover, during these periods, rats in the Cue Relevant group displayed significantly more active behaviors than did rats in the extinction or Cue Irrelevant condition (see Figure 1). Furthermore, during the first portion of the consummatory period, the rats in the anticipation group continued to show greater vigilance, exploration, and rearing than rats in the remaining two groups, but by the second sample within this period this activation had dissipated.

Interstitial Levels of CRH and GRP in Anticipation of a Palatable Snack

Figure 2 depicts the interstitial levels (presented as a percentage of baseline values) of CRH at the mPFC in response to the presentation of the 'anticipatory' cues and during the period associated with consumption (or not) of a palatable snack (Cue Relevant right and left, $n = 8$ and 10, respectively; Extinction right and left $n = 8$ and 6, respectively; Cue Irrelevant right and left, $n = 6$ and 6). The mixed measures ANOVA revealed that among animals with correctly positioned probes, CRH within the right mPFC varied as a function of the Treatment Condition \times Period interaction, $F_{2,38} = 4.01$, $p < 0.01$. The follow-up tests of the simple effects comprising this interaction indicated that in the Cue Relevant group and the Extinction condition, a significant rise of interstitial levels of CRH was evident during the 'anticipatory' period relative to their own baseline levels (see Figure 2). In both conditions, the levels of interstitial CRH remained elevated during the consummatory period, although a significant decline from the anticipatory period was apparent in the Cue Relevant group. Levels of CRH did not vary over samples in the Cue Irrelevant condition, and consequently, the interstitial CRH levels during the anticipatory and consummatory periods in

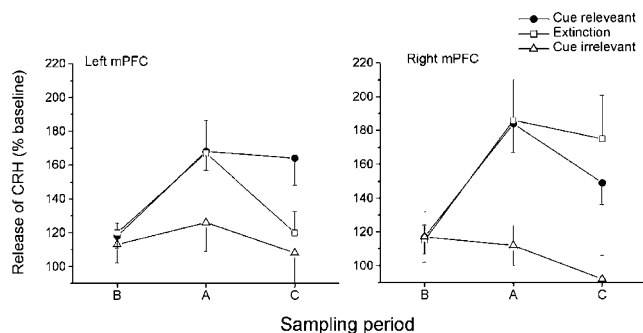


Figure 2 Effect of anticipatory cues and consumption of a palatable snack on the *in vivo* release of CRH in the left and right mPFC of rats in the Cue Relevant, Extinction, and Cue Irrelevant conditions. Scores represent the mean (\pm SEM) percentage change from the baseline period. Peptide changes are presented over the 40 min periods comprising the baseline period (B), the anticipation period (A), and after presentation of the palatable snack in the Cue Relevant group (C). Rats that were trained to 'anticipate' the presentation and consumption of palatable snack (Cue Relevant), similarly trained to anticipate a snack, but for a 2-week period prior to testing the snack presentation was discontinued (Extinction), or had the cues and the snack presented independent of one another (Cue Irrelevant). *Significantly greater than baseline, ($p < 0.01$); °significantly greater ($p < 0.01$) than the Cue Irrelevant group.

the Cue Relevant and Extinction groups significantly exceeded that of the Cue Irrelevant group.

Within the left mPFC, the interstitial CRH levels varied as a function of the Treatment Conditions \times Period \times Samples interaction, $F_{4,38} = 2.70$, $p = 0.04$. The profile of CRH changes was similar to that evident within the right mPFC, although these effects varied over samples within the anticipatory and consummatory phases. Specifically, the follow-up tests indicated that in both the Cue Relevant and Extinction conditions, CRH was elevated during the anticipation period relative to their own baselines. In the Cue Relevant group, the increase reached statistical significance during the second anticipatory sample, whereas in the extinction condition, the effect was significant during the first anticipatory sample. Whereas the rise was still evident in the Cue Relevant condition during the consummatory period, it fell to baseline levels in the Cue Irrelevant conditions. In contrast to the marked changes evident in the former conditions, CRH did not vary significantly over periods in the Cue Irrelevant group. Indeed, the follow-up comparisons indicated that while baseline CRH did not differ between the groups at baseline, the CRH levels in the Cue Relevant and Extinction groups exceeded that of the Cue Irrelevant group during the first anticipatory sample. Also, during the consummatory period, the CRH levels in the Cue relevant group exceeded that of the Cue irrelevant condition, whereas in the extinction condition this increase over the Cue Irrelevant condition was not sustained.

The interstitial levels (presented as a percentage of baseline values) of GRP at the mPFC in response to the presentation of the 'anticipatory' cues followed by the consumption of a palatable snack are shown in Figure 3 (Cue Relevant right and left, $n = 7$ and 10 , respectively; Extinction right and left $n = 8$ and 7 , respectively; Cue Irrelevant right and left, $n = 7$ /group). The interstitial levels of this peptide varied as a function of the Treatment

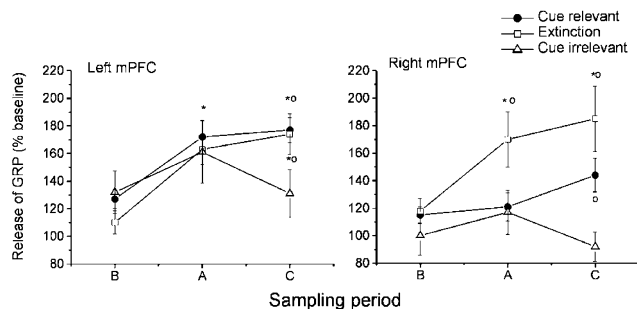


Figure 3 Effect of anticipatory cues and consumption of a palatable snack on the *in vivo* release of GRP in the left and right mPFC. Scores represent the mean (\pm SEM) percentage change from the baseline period. Peptide changes are presented over the 40 min periods comprising the baseline period (B), the anticipation period (A), and after presentation of the palatable snack in the Cue Relevant group (C). *Significantly greater than baseline ($p < 0.01$); °significantly greater ($p < 0.01$) than the Cue Irrelevant group.

Condition \times Period interactions, $F_{4,38} = 2.67$ and 2.60 ; p 's < 0.05 , for the right and left mPFC, respectively. The subsequent follow-up tests of the simple effects of this interaction confirmed that relative to baseline, in the Cue Relevant group, the anticipatory stimuli did not significantly increase interstitial GRP levels in the right mPFC, although there was an increase of GRP in the left hemisphere. Interestingly, the effects of the anticipatory stimuli on interstitial GRP levels were relatively marked in the Extinction group, where a significant rise of the peptide level was evident during the anticipation period, and continued to remain elevated (in both hemispheres) during the consummatory period. Relative to the levels evident in the Cue Irrelevant condition, in the extinction condition, the rise of GRP during the anticipatory and consummatory period was statistically significant in the right mPFC, but owing to a rise in the Cue Irrelevant group during the anticipatory period, between-group differences were noted during the consummatory period but not during the anticipatory period. In both hemispheres, the GRP levels in the Cue Relevant condition exceeded that of the Cue Irrelevant condition during the consummatory period.

Interstitial Levels of 5-HIAA and DA in Anticipation of a Palatable Snack

Figure 4 shows the interstitial levels of 5-HIAA at the mPFC (expressed as a percentage of baseline) in response to the presentation of the 'anticipatory' cues followed by the presentation (or not) of a palatable snack (Cue Relevant right and left, $n = 10$ and 10 , respectively; Extinction right and left $n = 8$ and 7 , respectively; Cue Irrelevant right and left, $n = 9$ and 10 , respectively). The accumulation of 5-HIAA in both the right and left mPFC varied as a function of the Treatment Condition \times Period interaction, $F_{4,44}$ and $F_{4,48} = 3.95$ and 3.79 , p 's < 0.01 . It appeared from the follow-up comparisons that in the Cue Relevant group, the interstitial accumulation of 5-HIAA increased during the anticipation period relative to baseline, but declined to baseline levels during the consummatory period. Accumulation of 5-HIAA was not apparent in either the Extinction or the Cue Relevant conditions. The follow-up comparisons

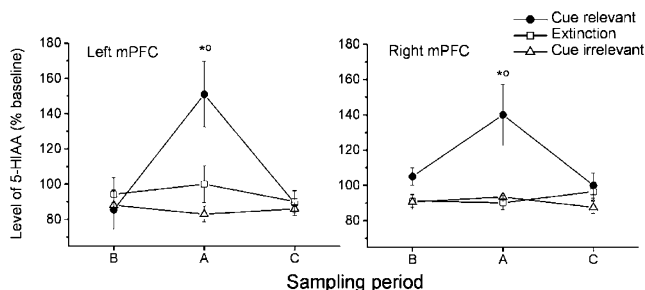


Figure 4 Effect of anticipatory cues and consumption of a palatable snack on the interstitial 5-HIAA levels at the left and right mPFC. Scores represent the mean (\pm SEM) percentage change from the baseline period. Peptide changes are presented over the 40 min periods comprising the baseline period (B), the anticipation period (A), and after presentation of the palatable snack in the Cue Relevant group (C). *Significantly greater than baseline ($p < 0.01$); °significantly greater ($p < 0.01$) than the Cue Irrelevant group.

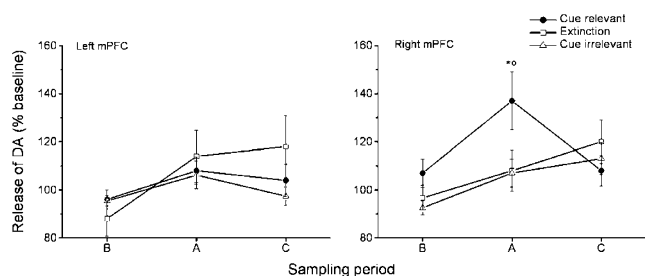


Figure 5 Effect of anticipatory cues and consumption of a palatable snack on the interstitial DA levels at the left and right mPFC. Scores represent the mean (\pm SEM) percentage change from the baseline period. Peptide changes are presented over the 40 min periods comprising the baseline period (B), the anticipation period (A), and after presentation of the palatable snack in the Cue Relevant group (C). *Significantly greater than baseline ($p < 0.01$); °significantly greater ($p < 0.05$) than the Cue Irrelevant group.

indicated that during the anticipation period, the 5-HIAA levels with the Cue Relevant group exceeded that evident in either of the other groups. The between-group differences were not evident either at baseline or during the consummatory phase.

The DA variations in the right and left mPFC as a function of the treatment conditions are shown in Figure 5 (Cue Relevant right and left, $n = 11$ and 10 , respectively; Extinction right and left $n = 8$ and 6 , respectively; Cue Irrelevant right and left, $n = 9$ and 6 , respectively). The interstitial levels of this amine within the right mPFC were found to vary as a function of the Treatment Condition \times Periods interaction, $F_{4,50} = 2.74$, $p = 0.03$. The follow-up comparisons confirmed that among rats in the Cue Relevant condition, the presentation of the anticipatory cues resulted in a significant increase of interstitial DA levels relative to baseline, and then declined to baseline levels during the consummatory period. In neither the extinction nor the Cue Irrelevant conditions was interstitial DA found to increase during the anticipation or consummatory periods. As a result, the DA levels during the anticipation period in the Cue Relevant condition significantly exceeded that in the former two conditions.

In contrast to the interstitial DA changes seen in the right mPFC, within the left mPFC, the levels of DA did not vary as a function of the Treatment or condition or the interactions involving this variable. There was a small rise of interstitial DA in the anticipation period (16%), but this increase was not different from that in the Extinction (15%) or Cue Irrelevant conditions (6%).

DISCUSSION

There have been several reports indicating that appetitive and aversive events may influence DA and 5-HT release in the nucleus accumbens and/or mPFC (eg Richardson and Gratton, 1996, 1998; Sullivan and Gratton, 1998). Also, food-related stimuli and changes of expected reward outcomes were found to promote DA changes within the nucleus accumbens (Blackburn *et al*, 1989; Richardson and Gratton, 1998). While food-related cues were reported to influence 5-HT release, less is known about the influence of anticipatory stimuli on 5-HT changes in limbic regions. Likewise, while CRH within the amygdala has been implicated in appetitive as well as aversive processes (Merali *et al*, 1998), it is uncertain whether changes of CRH are provoked in the mPFC in response to appetitive stimuli or cues that predict reward. Finally, GRP, which acts as a satiety peptide, has been shown to vary in response to food ingestion (Merali *et al*, 2002), but again, little is known about the GRP response to anticipatory stimuli. The present investigation assessed the influence of ingestion and anticipatory stimuli on neurochemical functioning within the mPFC, specifically whether such effects were unique to particular transmitters (eg variations of GRP, as a satiety peptide, would be aligned with ingestive processes, while DA and 5-HT changes would be evident in response to anticipatory cues).

Exposure to cues that had been paired with access to a palatable snack evoked behavioral signs of arousal. In contrast, when extinction training followed the same initial experience, active behaviors (exploration and vigilance) were no longer elicited by anticipatory stimuli. Instead, rats behaved like animals that had received irrelevant cue training, essentially being inactive most of the time. Thus, the presentation of the 'anticipatory' or incentive stimuli effectively elicited behavioral changes provided that alternative learning (extinction) had not taken place. Parenthetically, in animals with a restricted daily feeding schedule, increased activity is apparent approximately 1–3 h prior to feeding ('feeding anticipatory activity') (Aragona *et al*, 2002; Mistlberger and Marchant, 1999; Ono *et al*, 1996), a behavior that is attenuated by treatment with a 5-HT receptor antagonist (Ono *et al*, 1996; Shibata *et al*, 1995; Persons *et al*, 1993). In the present investigation, the increased activity in the Cue Relevant rats was only evident upon presentation of the anticipatory cues, thus the behavioral and neurochemical changes likely reflected the effects of incentive salience rather than a time-dependent general arousal. However, the possibility cannot be dismissed that the behavioral excitation in the Cue Relevant animals stemmed from frustration, resulting from the long delay (40 min) between cue presentation and animals actually receiving the snack (Dunn and Berridge, 1990; Heinrichs *et al*, 1995).

CRH has frequently been tied to motivational processes, but the influence of this peptide has typically been considered in the context of stressor effects. In this regard, stressor-provoked CRH variations have been observed at hypothalamic nuclei, the locus coeruleus, central amygdala, bed nucleus of the stria terminalis, as well as the mPFC (Bittencourt and Sawchenko, 2000; Curtis *et al*, 1995; Davis, 1998; Lavicky and Dunn, 1993). Although CRH receptors are widely distributed throughout the brain, including sites not typically associated with stressors (Potter *et al*, 1992; Bittencourt and Sawchenko, 2000), relatively little is known with regard to CRH involvement in behavioral processes such as those related to reward or anticipation of reward (Macey *et al*, 2000). Nevertheless, it seems that in addition to being activated in response to stressors, CRH release at the central amygdala may also be elicited by ingestion of a palatable snack (Merali *et al*, 1998). In the present investigation, it was further shown that in response to food-related anticipatory stimuli (Cue Relevant condition), CRH release increased within the mPFC. Interestingly, among rats that had been exposed to the extinction procedure, increased CRH was still evident upon presentation of anticipatory cues. The fact that this occurred, despite the absence of behavioral arousal, is consistent with the view that these cues maintain incentive value, but the extinction procedure resulted in other behavioral responses being adopted (Nader, 2003; Milad and Quirk, 2002).

Although the satiety effects of GRP (and related peptides, for example, bombesin) have been extensively assessed (Babcock *et al*, 1985; Gibbs, 1985; Gibbs *et al*, 1979; Gibbs and Smith, 1988; Merali *et al*, 1999, 2002), there is a paucity of information concerning the involvement of this peptide in relation to reward or anticipatory mechanisms. In the current investigation, the Cue Relevant and Extinction groups displayed comparable increases of GRP in response to the anticipatory cues; but, in the Cue Relevant group, this increase was most pronounced once rats engaged in actual food ingestion. Curiously, in the Extinction condition, the increase was significant during the 'anticipatory' period. While interesting, these findings are difficult to explain. It has been reported that in rats with stimulating electrodes located in the lateral hypothalamus, intraperitoneal administration of bombesin increased the threshold for stimulation-induced feeding, but did not influence intracranial self-stimulation. Thus, it appeared that the effect of this peptide on stimulation-induced feeding was similar to that of normal feeding (Bushnik *et al*, 1999). Yet, the finding that GRP release was more pronounced in the Extinction than in the Cue Relevant condition suggests that this peptide is affected by processes unrelated to feeding or satiety *per se*. Given that GRP is released not only in response to food intake but also in response to stressor application (Kent *et al*, 1998; Merali *et al*, 1998), the possibility exists that the frustration/distress associated with the snack being withheld (in the Extinction condition) was responsible for the increased release of the peptide, even though animals did not show behavioral signs of distress and generally seemed unresponsive in the presence of the cue that had previously been associated with snack delivery.

It has been reported that DA release at the mPFC and/or the nucleus accumbens was increased among rats responding for food reward (Hernandez and Hoebel, 1988) or in

response to relatively novel foods (Ahn and Phillips, 1999; Bassareo and Di Chiara, 1997; Feenstra, 2000; Feenstra and Botterbom, 1996). As DA neuronal activity was also affected by perturbations of expected reward (delaying reward presentation or reducing reward duration), it was suggested that DA neuronal activity operates to influence the incentive value of the food reward (Richardson and Gratton, 1996, 1998). Likewise, the view was offered that the mPFC, in conjunction with CeA, operates to influence the sensory incentive properties of food (Ahn and Phillips, 1999, 2002). Within the present investigation, anticipatory cues were indeed found to affect DA efflux, but this outcome was apparent only within the right mPFC. Within the left mPFC, there was a small rise of DA release in response to anticipatory cues, but this effect did not approach statistical significance.

Several studies have highlighted left/right hemispheric functional asymmetries involving mPFC DA activity following exposure to stressors (Sullivan and Gratton, 2002, 1998; Nielsen *et al*, 1999; Brake *et al*, 2000). It was indeed suggested that, depending on the type of stressor involved, the right mPFC may have a specialized role in the integration of emotional and physiological responses in aversive situations (Sullivan and Gratton, 1998; Brake *et al*, 2000). Yet, using positron emission tomography, it was shown that stimuli associated with cocaine (which elicit cocaine craving) provoked activation of several sites within the right hemisphere (eg dorsolateral prefrontal cortex) and deactivation of the left frontopolar and mPFC. Thus, an overall bias towards the right hemisphere activation would be elicited by these cues (Bonson *et al*, 2002). Given that DA variations in the present investigation were only apparent within the right mPFC suggests that a similar asymmetry also occurs with respect to anticipatory response related to food reward. Importantly, however, the elevated DA release was not evident during the consummatory period. Thus, the possibility exists that the right mPFC DA elevations in response to anticipatory cues may have reflected adverse effects (eg frustration) related to the lengthy period of anticipation (40 min) prior to actual food delivery.

While it is clear that asymmetry existed with respect to the effects of anticipatory stimuli on DA release, this does not necessarily suggest that DA changes are not elicited in the left mPFC. At this juncture, we can only conclude that the right mPFC is more sensitive than the left in response to anticipatory cues. Moreover, actual food ingestion did not affect DA release in either hemisphere. It is uncertain whether the absence of more profound DA variations within the left mPFC reflected the fact that animals had been well trained to anticipate the food, hence leading to devaluation of its novelty, or whether it stemmed from relatively transient changes of DA being obfuscated by the fairly long (20 min) sampling periods required for detection of the amine. Further, given that DA changes in the nucleus accumbens and striatum associated with ingestion of a nutritive meal were previously found not to be apparent in response to a palatable non-nutritive saccharin solution (Blackburn *et al*, 1989), it is similarly possible that such a factor may have limited the mPFC DA changes associated with the palatable snack in rats that had not been food deprived. Also, accumbal DA release was shown to occur in conjunction with the emission of operant responses, rather

than the actual consumption of food reward (Salamone *et al*, 1994); thus, it ought to be considered that DA variations within the left mPFC might be associated with actual responding for appetitive reward, rather than anticipatory factors or ingestion *per se*.

In addition to DA, hypothalamic 5-HT has been implicated in the regulation of food intake (Simansky, 1998). Moreover, the release of this amine is increased in response to food odors or cues that had been paired with food (Hoebel *et al*, 1989; Schwartz *et al*, 1990), suggesting that this monoamine may be important in the anticipation associated with food intake. Given that 5-HT receptors modulate cortical DA function in the mPFC (Pehk *et al*, 2001), and 5-HT has been implicated as a contributing factor in the rewarding effects of psychostimulants (Brown and Molliver, 2000; Kaga *et al*, 2001), the activity of this amine might have been expected to be influenced by stimuli with incentive properties. Indeed, in the present investigation, 'anticipatory' stimuli increased 5-HIAA accumulation in the Cue Relevant group, just as these stimuli increased CRH release. However, unlike the CRH changes, in the Extinction condition, there was no evidence of elevated 5-HIAA or DA in response to the anticipatory stimuli. Thus, greater selectivity was evident with respect to change in these amines than there were with respect to CRH release. To be sure, accumulation of the 5-HT metabolite, 5-HIAA, does not necessarily reflect the release of 5-HT. Among other things, treatments may affect uptake processes so that the 5-HT and 5-HIAA levels under varied conditions differ from one another. Yet, the fact that the 5-HIAA levels were selectively increased in the Cue Relevant group is consistent with the position that relevant stimuli promoted 5-HT release. That a similar outcome was not apparent in the other two groups cannot necessarily be taken to suggest that increased 5-HT release was not elicited by the treatments, although the data would imply that any such effect is more pronounced in the Cue Relevant condition.

Cues associated with a palatable snack provoked increased activity of CRH, 5-HT, and DA neurons; however, the functional significance of these neurochemical changes was likely different from one another. Specifically, once animals had learned that the cues no longer predicted snack presentation, the increased CRH release persisted, whereas that of 5-HT and DA did not. It is conceivable that peptidergic (CRH and possibly GRP) systems serve to allocate salience and/or incentive reward value to biologically significant stimuli or reflect the emotional response to anticipatory stimuli, even after animals have undergone extinction. As indicated earlier, extinction may not eliminate the potential salience of the cues, but rather promotes the adoption of other, more appropriate, responses. In contrast, 5-HT neuronal activity and that of DA within the right mPFC may be more closely aligned with the cognitive appraisal associated with predictor stimuli, such that the increased neuronal activity would be limited to conditions where reward was eminent.

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