Motivational deficits in individuals at-risk for psychosis and across the course of schizophrenia

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A R T I C L E   I N F O

Article history:
Received 31 January 2014
Received in revised form 18 June 2014
Accepted 19 June 2014
Available online 5 July 2014

Keywords:
Schizophrenia
Motivation
Avolition
Anticipatory Pleasure
Psychosis
Reward
Duration of illness

A B S T R A C T

Motivational impairment is a critical factor that contributes to functional disability in schizophrenia and under- mines an individual’s ability to engage in and adhere to effective treatment. However, little is known about the developmental trajectory of deficits in motivation and whether these deficits are present prior to the onset of psychosis. We assessed several components of motivation including anticipatory versus consummatory pleasure (using the Temporal Experience of Pleasure Scale (TEPS)), and behavioral drive, behavioral inhibition, and reward responsivity (using the Behavioral Inhibition Scale/Behavioral Activation Scale (BIS/BAS)). A total of 234 participants completed study measures, including 60 clinical high risk (CHR) participants, 60 recent-onset schizophrenia participants (RO), 78 chronic schizophrenia participants (SZ) and 29 healthy controls (HC) age matched to the CHR group. CHR participants endorsed greater deficits in anticipatory pleasure and reward responsivity, relative to HC comparison participants and individuals diagnosed with schizophrenia. Motivational deficits were not more pronounced over the course of illness. Depressed mood was uniquely associated with im- pairments in motivation in the CHR sample, but not the schizophrenia participants. The results suggest that CHR individuals experience multiple contributors to impaired motivation, and thus multiple leverage points for treatment.

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1. Introduction

Dating back to the earliest descriptions of schizophrenia by Kraepelin and Bleuler, amotivation/avolition was observed to be central to the phenomenology and course of schizophrenia (McGlashan, 2011). Amotivation and negative symptoms more broadly are resistant to cur-

Cohen and Minor, 2010; Foussias and Remington, 2010), as well as difficulty maintaining cognitive representation of rewarding experiences and redirecting behavior back to rewarding experiences (Barch and Dowd, 2010).

While much of the research on motivation in schizophrenia has fo-

http://dx.doi.org/10.1016/j.schres.2014.06.024
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for negative symptoms and motivation specifically. In the first study, Scholten et al. (2006) found that individuals with schizophrenia were more sensitive to threat than healthy controls, but no differences were detected between patients and controls in the behavioral activation system. In a more recent study (Engel et al., 2013), the BAS system was found to be negatively associated with more severe negative symptoms, suggesting heterogeneity within schizophrenia samples. Thus far, most studies on motivation and reward processing in schizophrenia have been conducted on individuals who have been persistently ill for most of their adult lives. It is therefore unknown whether motivational deficits worsen over the course of schizophrenia. The stability of negative symptoms generally, however, has been examined and while it appears they might be stable in severity across the course of psychosis, some studies suggest that as the duration of untreated psychosis increases, individuals with schizophrenia experience worsening negative symptoms over time (Chang et al., 2013). Although motivational deficits are often considered within the context of negative symptoms and reward processing, the presence of mood and anxiety symptoms is also linked to motivational capacity in schizophrenia. In particular, depressed mood is associated with decreased hedonic capacity while anxiety symptom severity is associated with greater threat sensitivity/avoidance behavior (Barch et al., 2008). These studies raise questions about the degree of behavioral activation and avoidance within schizophrenia samples that vary in the degree of negative symptom severity, as well as other factors, such as the duration of illness, and the presence of comorbid depressive and anxiety symptoms. In the current study, we examined negative symptom severity, mood and anxiety symptoms, and motivational deficits across the course of schizophrenia, using a cross-sectional design.

The primary aim of this study was to examine several behavioral components of motivation (wanting, liking, approach and avoidance) in individuals at various stages of experiencing psychosis: those at clinical high risk (CHR) of developing a psychotic disorder, those within the first 5 years of onset of schizophrenia (Recent Onset; RO), and those with persistent schizophrenia or schizoaffective disorder (SZ). We tested the following hypotheses: 1) Individuals with persistent schizophrenia will demonstrate greater motivational impairments than those at-risk or with a recent onset of schizophrenia, and 2) mood and anxiety symptoms will influence the degree of motivational impairments at all stages of illness, such that more severe anxiety and depression will be positively correlated with avoidance and negatively correlated with wanting, liking, and approach motivation in all participant groups.

2. Methods

2.1. Participants

The study included 234 participants: 60 clinical high risk (CHR), 60 recent-onset schizophrenia (RO), 78 persistent schizophrenia (SZ) participants and 29 healthy controls (HC) who were recruited for randomized controlled trials of cognitive training (ClinicalTrials.gov NCT00655239, NCT00694889, and NCT00312962). The HC participants were age-matched to the CHR participants. Participants in the HC, CHR and RO groups were drawn from two research programs at the University of California, San Francisco (UCSF) and University of California at Davis (UCD) and persistent SZ participants from a research program at the San Francisco VA Medical Center (SFVAMC). Patient participants were recruited from community mental health centers, outpatient clinics, local schools and universities, and HC participants were recruited via advertisement. CHR status was ascertained using the Structured Interview for Prodromal Syndromes (SIPS version 4.0 (Miller et al., 2002). All CHR participants met one of the following prodromal syndromes on the SIPS/SOPS: 1) the presence of attenuated positive, psychotic symptoms, occurring at least weekly with onset or worsening in the past year, 2) brief intermittent psychotic symptoms, which must have begun in the past three months, or 3) a 30% decline in GAF score over the past year, plus either a diagnosis of schizotypal personality disorder or a first-degree relative with a psychotic disorder. Recent onset participants were included based on a diagnosis of schizophrenia, schizophreniform, or schizoaffective disorder with an onset within the past five years as determined by the Structured Clinical Interview for the DSM-IV TR Axis I disorder interview (SCID-I; First, 1996). Onset of illness was defined by the date diagnostic criteria were first met, as assessed by the SCID. Persistent schizophrenia participants were diagnosed using the SCID-I. Healthy controls did not meet DSM-IV criteria for an Axis I psychiatric disorder as determined by the SCID-I or meet criteria for a prodromal syndrome, and had no first-degree relatives with psychosis, based on participant and collateral informant reports during the SIPS interview. CHR, RO and SZ participants were clinically stable at the time of testing (no hospitalization within the past 3 months and stable dose of medication over the past month), as per the requirements for the parent cognitive training study. Other inclusion criteria included: 1) good general physical health; 2) fluent and proficient in English; 3) IQ > 70 (WASI, 1st edition, 2-subtest version: Vocabulary and Matric Reasoning); 4) no neurological disorder; and 5) no substance dependence or significant use that would interfere with study participation.

2.2. Procedure

Advanced graduate students, predoctoral interns, postdoctoral fellows, and trained bachelor-level research assistants administered the measures described below in the context of a larger battery of cognitive and clinical assessments. All participants gave written informed consent or assent for the study and were compensated for their participation in all assessments. Parental informed consent for minors was also obtained. After an intake evaluation that determined study eligibility, all participants underwent a structured diagnostic clinical interview and completed self-report measures of motivation and clinician ratings of symptom severity. Only baseline, cross-sectional data were included in this study.

2.3. Measures

We assessed several components of motivation including negative symptom severity and mood and anxiety using the Positive and Negative Syndrome Scale (PANSS; Kay et al., 1987), anticipatory (wanting) versus consummatory (liking) pleasure using the Temporal Experience of Pleasure Scale (TEPS; Gard et al., 2006), and behavioral drive, behavioral inhibition, and reward responsivity using the Behavioral Inhibition Scale/Behavioral Activation Scale (BIS/BAS; Carver and White, 1994). The TEPS includes eighteen items and are rated on a scale of 1 (very false for me) to 6 (very true for me). The BIS/BAS measure is comprised of twenty-four items rated on scale of 1 (very true for me) to 4 (very false for me), of which seven items represent the BIS. The BAS scales were designed to measure approach motivation traits while the BIS scale was designed to measure aversive motivation traits. Examples of TEPS and BIS/BAS items are in Table 1. The PANSS is clinician-administered, while the TEPS and BIS/BAS are self-report measures.

2.4. Data analytic plan

First, data were inspected for normality and outliers were winsorized at a level of 95%. Less than 5% of the data on these measures, within each group, were adjusted. One-way ANOVAs and chi-square tests were conducted to test for demographic differences. To better understand the relationship between the TEPS and BIS/BAS scales, we used Pearson correlation analyses. BIS/BAS scores were reverse coded, such that higher ratings represented greater degrees of approach motivation. To test hypothesis 1, a series of one-way ANOVAs were used to compare the mean differences in anticipatory and consummatory pleasure and approach and aversive motivation between groups. In order to test Hypothesis 2, mood and anxiety symptoms were examined
in relation to motivation, using Pearson correlation analyses. If significant, the PANSS mood and/or anxiety ratings were included as covariates in the main analyses (ANCOVAs). Post hoc analyses were conducted using Tukey tests.

3. Results

The demographic characteristics of the study participants are included in Table 2. The HC participants were matched to the CHR subjects by age, gender, education, and IQ. The RO and SZ participants included significantly more males and a lower IQ than the HC and CHR samples. The SZ sample was significantly more educated than the HC and CHR participants, likely due to age differences. Negative symptoms were more severe in the schizophrenia samples (RO and SZ) than the CHR sample and there was no significant difference in negative symptom severity between RO and SZ. Demographic characteristics (age, gender, education, and IQ) were not associated with the TEPS and BIS/BAS ratings in each sample.

The interrelationships between TEPS and BIS/BAS scales (Table 3) demonstrated a strong, positive relationship between reward responsivity and the anticipatory pleasure scale in all samples. Reward responsivity demonstrated a strong, positive relationship between reward responsivity (approach motivation traits) and behavioral inhibition in each sample; however, anxiety was found to be strongly related to behavioral inhibition in the RO sample, while only related to consummatory pleasure in HCs.

Due to their potential relationships with motivation, depression and anxiety severity scores were examined in each sample. Depressed mood was strongly, negatively correlated with anticipatory pleasure in CHR participants \((r = -0.430, p = .002)\), but not in the RO or SZ samples. Interestingly, there was no significant difference in the severity of depressed mood between CHR and RO; however, the RO and SZ participants exhibited more severe negative symptom severity, relative to the CHR participants. Depressed mood was unrelated to behavioral approach (drive or reward responsivity) or behavioral inhibition in each sample; however, anxiety was found to be strongly related to behavioral inhibition in the CHR \((r = .564, p < .001)\), RO \((r = .232, p = .03)\), and SZ \((r = .377, p = .001)\) samples. Due to the significant relationship between depressed mood and anticipatory pleasure in the CHR group, depressed mood was included as a covariate in the main analysis comparing group differences in anticipatory pleasure. Anxiety severity was also included in the main analysis comparing group differences in behavioral inhibition.

The overall between-group differences in anticipatory and consummatory pleasure were significant \((F(3,224) = 7.33; p = .001; \text{Fig. } 1)\). This result remained significant when depressed mood was entered as a covariate. Post hoc comparisons using the Tukey test indicated that this results was driven by CHR participants who reported significantly less anticipatory \((M = 3.61, SD = .73)\) and consummatory pleasure \((M = 3.84, SD = 1.1)\), than HC (anticipatory pleasure: \(M = 4.48, SD = .73\); consummatory pleasure: \(M = 4.42, SD = .91\) ), RO (anticipatory pleasure: \(M = 4.17, SD = .89\); consummatory pleasure: \(M = 4.39, SD = .80\)), and SZ participants (anticipatory pleasure: \(M = 4.11, SD = .93\); consummatory pleasure: \(M = 4.31, SD = .11\)). The RO and SZ samples endorsed intact anticipatory and consummatory pleasure, relative to the young HC sample.

The within-group differences in anticipatory versus consummatory pleasure were examined in each of the samples (Fig. 2). No significant difference in anticipatory versus consummatory pleasure was observed in HCs \((t = .47, p = .64)\). In both the RO and SZ samples, their anticipatory pleasure was reported to be significantly lower than consummatory pleasure \((RO: t = -2.33, p = .02; SZ: t = 2.00, p = .05)\). In CHR participants, this difference was at trend level \((t = -1.85, p = .07)\). The results remained significant for the RO and SZ groups when depressed mood was entered as a covariate.

Between-group differences in behavioral drive and reward responsivity (approach motivation traits) were tested (Fig. 3). The overall differences in behavioral drive and reward responsivity were significantly different between the groups (behavioral drive: \(F(3, 230) = 7.46, p = .001\); reward responsivity: \(F(3, 230) = 9.17, p = .001\)). Consistent with the directionality of the anticipatory and consummatory pleasure data, CHR participants endorsed lower behavioral drive \((M = 2.37, SD = .73)\), relative to RO \((M = 2.83, SD = .63)\) and SZ \((M = 2.81, SD = .68)\) participants, and lower reward responsivity \((M = 3.03, SD = .58)\) relative to the HC, \((M = 3.40, SD = .35)\) RO \((M = 3.40, SD = .41)\) or SZ \((M = 3.42, SD = .49)\) participants. There was no difference in the approach motivation traits between any of

<table>
<thead>
<tr>
<th>TEPS-Anticipatory pleasure</th>
<th>TEPS-Consummatory pleasure</th>
<th>Behavioral inhibition</th>
<th>Behavioral drive</th>
<th>Reward responsivity</th>
</tr>
</thead>
<tbody>
<tr>
<td>When something exciting is coming up in my life, I really look forward to it. I have noticed that looking forward to a pleasurable experience is in itself pleasurable</td>
<td>I have enjoyed experiences like taking deep breaths of fresh air when I walk outside. The smell of freshly cut grass (or the outdoors) has been enjoyable to me</td>
<td>I feel pretty worried or upset when I think or know somebody is angry at me. I worry about making mistakes</td>
<td>When I want something, I usually go all-out to get it. I go out of my way to get things I want</td>
<td>When I get something I want, I feel excited and energized. When good things happen to me, it affects me strongly</td>
</tr>
</tbody>
</table>

### Table 2: Demographic characteristics of study participants.

<table>
<thead>
<tr>
<th>Healthy controls (n = 29)</th>
<th>Clinical high risk (n = 60)</th>
<th>Recent onset (n = 67)</th>
<th>Schizophrenia (n = 78)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>17.6 (3.7)</td>
<td>18.6 (4.6)</td>
<td>21.7 (4.5) *</td>
</tr>
<tr>
<td>Male (%)</td>
<td>48%</td>
<td>52%</td>
<td>70%</td>
</tr>
<tr>
<td>Caucasian (%)</td>
<td>63%</td>
<td>57%</td>
<td>60%</td>
</tr>
<tr>
<td>Subject education (mean years, SD)</td>
<td>11.4 (2.9)</td>
<td>11.5 (3.1)</td>
<td>12.8 (1.9)</td>
</tr>
<tr>
<td>WASI-IQ</td>
<td>110.9 (13.5)</td>
<td>105.6 (16.0)</td>
<td>102.8 (12.8) **</td>
</tr>
<tr>
<td>Months of illness (mean, SD)</td>
<td>–</td>
<td>–</td>
<td>21.9 (2.5) **</td>
</tr>
<tr>
<td>Negative symptoms (mean, SD)</td>
<td>–</td>
<td>4.68 (50) **</td>
<td>17.89 (6.1) **</td>
</tr>
<tr>
<td>Depression (mean, SD)</td>
<td>3.18 (1.8) *</td>
<td>2.50 (1.6) *</td>
<td>3.14 (1.6) *</td>
</tr>
<tr>
<td>Anxiety (mean, SD)</td>
<td>3.36 (1.3)</td>
<td>2.85 (1.5)</td>
<td>3.26 (1.3)</td>
</tr>
</tbody>
</table>

Note. Recent onset and schizophrenia participants were significantly older, were more represented by males, were more educated, and had a lower IQ than healthy control and clinical high risk participants. Recent onset and schizophrenia participants had more severe negative symptoms and less severe depressive symptoms than clinical high risk participants.

* \( p < .05 \).

** \( p < .01 \).
the other paired participant groups. Depression and anxiety were not entered as covariates since these were unrelated to drive and reward responsivity in each of the samples.

Behavioral inhibition was significantly different between the groups (F(3,230) = 3.39; p = .02). Consistent with the pattern in the behavioral approach data, CHR participants endorsed greater behavioral inhibition (M = 3.04, SD = .66) than the HCs (M = 2.68, SD = .42), but no differences were observed between any other paired groups. The results remained the same when anxiety was entered as a covariate.

There were no significant differences between the RO and SZ participants on any of the motivational items, suggesting no decline in anticipatory pleasure, reward responsivity, drive or inhibition across the duration of illness. Further, the correlations between months of illness and anticipatory and consummatory pleasure, behavioral approach and inhibition were not significant (all r’s < .100).

4. Discussion

This is the first study, to our knowledge, to investigate motivational deficits across individuals with psychosis of varying illness duration. Notably, CHR participants endorsed more severe deficits in anticipatory pleasure, consummatory pleasure, and reward responsivity, relative to age-matched healthy comparison subjects, and, contrary to our hypotheses, also relative to individuals already diagnosed with early-course and later-course schizophrenia. CHR participants also showed lower behavioral drive relative to individuals with early and later-course schizophrenia. This is surprising, considering that overall negative symptoms were more pronounced in the established schizophrenia samples. Our hypothesis that motivational deficits would be more severe in individuals living with schizophrenia for a greater duration was also not supported. CHR participants endorsed greater motivational deficits across a range of motivational factors compared to early illness and chronic schizophrenia participants, even after controlling for depressed mood.

The degree and extent of motivational impairments endorsed by CHR participants exemplifies the multifarious sequelae of the CHR syndrome. Individuals who are at imminent risk for psychosis are phenotypically complex. Many CHR individuals experience a multitude of problems, including attenuated psychotic symptoms, depressed mood, anxiety, social withdrawal, and functional deficits, all of which might reflect the behavioral correlates of a reward system that is impaired. Further, the degree to which the reward system appears to be impaired has predictive validity for this population. Less severe negative symptoms and depressed mood in CHR individuals significantly increases the likelihood of recovering from an at-risk state and experiencing improved functional outcomes (Addington et al., 2011; Schlosser et al., 2012). Furthermore, it is possible that motivational deficits in CHR individuals who go on to develop psychosis might be a precursor to the development of negative symptoms, if they are not yet present for an individual. Taken together, this suggests that motivational deficits should be a target of vigorous preemptive treatment.

The wanting vs liking discrepancy that has been previously reported in the literature was observed among the RO and SZ samples and was at trend level in the CHR group, suggesting that a characteristic feature of schizophrenia may indeed be an impairment in the anticipation of rewarding stimuli and experiences. Also consistent with the literature, RO and SZ participants reported intact consummatory pleasure. However, CHR participants reported significantly less consummatory and

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**Table 3**

<table>
<thead>
<tr>
<th></th>
<th>Behavioral Inhibition</th>
<th>Reward Responsivity</th>
<th>Behavioral Drive</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Healthy control</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anticipatory pleasure</td>
<td>.254 ± .073</td>
<td>.548** ± .026</td>
<td>.347 ± .009</td>
</tr>
<tr>
<td>Consummatory pleasure</td>
<td>-.174 ± .036</td>
<td>.379* ± .012</td>
<td>.461** ± .004</td>
</tr>
<tr>
<td><strong>Clinical high risk</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anticipatory pleasure</td>
<td>.022 ± .005</td>
<td>.459** ± .012</td>
<td>.128 ± .003</td>
</tr>
<tr>
<td>Consummatory pleasure</td>
<td>-.149 ± .029</td>
<td>.081 ± .004</td>
<td>.014 ± .001</td>
</tr>
<tr>
<td><strong>Recent onset</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anticipatory pleasure</td>
<td>.050 ± .036</td>
<td>.591** ± .012</td>
<td>.389** ± .009</td>
</tr>
<tr>
<td>Consummatory pleasure</td>
<td>.022 ± .005</td>
<td>.477** ± .012</td>
<td>.273* ± .004</td>
</tr>
<tr>
<td><strong>Schizophrenia</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anticipatory pleasure</td>
<td>.263* ± .014</td>
<td>.424** ± .012</td>
<td>.181 ± .004</td>
</tr>
<tr>
<td>Consummatory pleasure</td>
<td>.185 ± .014</td>
<td>.424** ± .012</td>
<td>.151 ± .004</td>
</tr>
</tbody>
</table>

*p < .05.

**p < .01.

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**Fig. 1.** Between group differences of anticipatory and consummatory pleasure.

**Fig. 2.** Within-group differences in anticipatory and consummatory pleasure.

**Fig. 3.** Between group differences in behavioral drive and reward responsivity.
Role of funding source

This research was supported by the following grants: NIMH K23 MH097795-01 (Schlosser); UCSF CTSA S01 UL1 TR000004; Schlosser); NIMH R34 MH100399 (Schlosser, Vinogradov) and T32 MH089820 (Ford). The funding sources had no further role in study design; in the collection, analysis and interpretation of data; in the writing of the report; and in the decision to submit the paper for publication.

Contributors

We confirm that the manuscript has been read and approved by all named authors and that there are no other persons who satisfied the criteria for authorship but are not listed. We further confirm that the order of authors listed in the manuscript has been approved by all of us.

Conflict of interest

We wish to confirm that there are no known conflicts of interest associated with this publication and there has been no significant financial support for this work that could have influenced its outcome.

Acknowledgments

We would like to thank the participants and their families for contributing to this research.

References


Harre, R., Parrott, W.G., 1996. The Emotions: Social, Cultural and Biological Dimensions. SAGE.


