Emotion-Regulation in Psychosis: Patients with Psychotic Disorders Apply Reappraisal Successfully

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Abstract

Background. Emotion-regulation is assumed to be impaired in psychosis: patients with psychotic disorders (PD) self-report to use reappraisal less frequently than healthy controls (HC), but it is uncertain whether they are also less successful in applying reappraisal. Moreover, it has not been investigated whether the habitual use of reappraisal is associated with the ability to use reappraisal successfully.

Methods. To address these questions, PD (n = 60), a clinical control group of patients with anxiety disorders (AD; n = 40) and HC (n = 40) completed questionnaires on habitual emotion-regulation and alexithymia, performed tests on neurocognitive functioning and jumping-to-conclusion bias, and were subjected to an emotion-regulation paradigm, in which they were asked to down-regulate induced anxiety and sadness.

Results. No differences were found between PD and HC regarding the habitual use of reappraisal, but PD reported to use reappraisal more frequently than AD. All groups were able to down-regulate anxiety and sadness via reappraisal following instructions. The habitual use of reappraisal and the reduction of anxiety and sadness after instruction were uncorrelated.

Conclusion. These findings question the common assumption that PD use reappraisal less often or less successfully. At least under laboratory condition, PD do not appear to be impaired in reducing negative affect via reappraisal successfully.

Keywords. Psychosis · Schizophrenia · Emotion regulation · Reappraisal

High rates of depression and anxiety are found in patients with psychotic disorders (PD; meta-analyses: Achim et al. 2011; Li et al. 2020). Negative affect, such as depression and anxiety, is associated with lower quality of life (e.g. Gardsjord et al. 2016; Huppert et al. 2001) and is postulated to contribute to the formation and maintenance of psychotic symptoms (Garety et al. 2001; Hartley et al. 2013). Thus, it does not seem surprising that psychosis is also associated with deficits in emotion-regulation (Ludwig et al. 2019), which was originally defined as “extrinsic and intrinsic processes responsible for monitoring, evaluating, and modifying emotional reactions, especially their intensive and temporal features, to accomplish one’s goals” (Thompson 1994, pp.27–28) and has undergone numerous revisions and extensions since then (e.g. Gross 2015; Hofmann 2014; Hofmann et al. 2012; Koole 2009; Tamir et al. 2015). In this study we focused on one specific emotion-regulation strategy, namely reappraisal.

Reappraisal in the context of emotion regulation is commonly defined as cognitively transforming a situation in order to change its emotional impact (Gross 1998). For example, being passed by a colleague on the street without being greeted could spontaneously be interpreted as being disliked by that person, which may cause a person to feel sad or angry on a subjective level and may also be associated with higher levels of arousal, which may then be evaluated as threatening (Freeman et al. 2002). However, being able to then consider alternative explanations, such as the colleague being in a hurry, could buffer the initial emotional reaction. Reappraisal has been found to be effective on a short-term basis in the reduction of negative affect in experimental designs and the habitual use of reappraisal has been found to be functional in terms of being associated with adaptive long-term outcomes, such as healthier patterns of affect, social functioning, and well-being in healthy adults.
(John & Gross 2004; McRae et al. 2012a). An effect of reappraisal has also been demonstrated on physiological indicators of arousal (e.g. Hofmann et al. 2009; Jackson et al. 2000; McRae et al. 2012b; Ray et al. 2010).

Over and beyond the observation that negative affect is high in people with psychosis, there are several reasons to expect PD to have pronounced difficulties in using reappraisal successfully: First, the ability to identify and describe one’s emotions, which is considered a prerequisite of effective reappraisal (Barrett et al. 2001; Füstös et al. 2012; Van Rijn et al. 2011), has been found to be impaired in psychosis (alexithymia, O’Driscoll et al. 2014). Second, deficits in neurocognitive functioning could hamper the successful implementation of a cognitively demanding strategy such as reappraisal (Schmeichel and Tang 2015). Third, acute psychosis is associated with a tendency to jump to conclusions, i.e. making unwarranted decisions on the basis of little evidence (Dudley et al. 2015). This cognitive bias is also likely to antagonize attempts to generate and decide on helpful alternative explanations and therefore likely to impede reappraisal. Nevertheless, it remains to be shown whether these factors are indeed related to the ability to apply reappraisal successfully.

Previous research on emotion-regulation and psychosis has mostly focused on the self-reported habitual use of reappraisal. In these studies, participants are typically asked to report if or how often they use reappraisal. A meta-analysis of questionnaire-based studies found that PD report to use reappraisal habitually to a lesser extent compared to healthy control subjects (HC) with a small to moderate effect size (Ludwig et al. 2019), but studies comparing PD to clinical control groups are scarce. This is surprising, given that emotion-regulation deficits are prevalent in numerous other disorders, particularly in affective and anxiety disorders (Picó-Pérez et al. 2017). Thus, comparing PD to patients with these disorders is of interest in order to assess psychosis-specificity. Also, a problem with the types of questionnaires used is that they tend not to assess success but are restricted to the analysis of how frequently certain emotion-regulation strategies are used due to the way the questions are phrased and scaled (e.g. item from the Cognitive Emotion Regulation Questionnaire (CERQ): ‘I look for the positive sides to the matter’ with a scale ranging from ‘never’ to ‘always’ Garnefski et al. 2002). However, the amount of using reappraisal does not necessarily reveal information about the ability to successfully implement the strategy. To assess this ability, we need to measure the reduction in negative affect following the use of a strategy.

To date, only few studies have used experimental designs to examine the successful application of reappraisal in PD. These studies typically use emotion-eliciting stimuli, such as pictures, to induce emotions and subjects are asked to either apply certain emotion-regulation strategies (experimental conditions) or to just view (control condition) and to rate their subjective emotion intensity. Three such studies found no deficits in the ability to implement reappraisal successfully in PD (Grezellschak et al. 2015; Painter et al. 2019; Perry et al. 2012). Only one study found evidence for potential deficits in reappraisal success, but this study compared subclinical participants with high delusion-proneness to participants with low delusions-proneness (Westermann et al. 2014). Beyond the low number of available studies, a problem with those at hand is that they predominantly (1) comprise small samples that are not suited to detect small effects, (2) do not include more than one indicator of arousal, which is important as emotion regulation is assumed to have an impact on all levels of emotional responding (Koole, 2009) (3) do not include more than one emotion to examine emotion-specificity, even though a predictive value of affect on psychotic symptoms has predominantly been found for anxiety (Freeman et al. 2015; Krkovic et al. 2019; Lincoln et al. 2009; Oliver et al. 2012; Thewissen et al. 2011) and theoretical models have stressed anxiety as being particularly relevant to delusion formation (e.g. Freeman et al. 2002) and (4) do not include clinical control groups to examine psychosis-specificity. Thus, we are still unable to answer the question of whether psychosis is associated with a deficit in the ability to apply reappraisal successfully.

Finally, the separate strands of research on the successful use of reappraisal in experimental paradigms and the habitual use of reappraisal reported in questionnaires have hardly been related to each other so far. It seems plausible that if an individual is not able to reduce negative affect via reappraisal, this individual would have little incentive to apply that strategy habitually. Vice versa, a strategy that is seldom used gets little practice, which could impede its effectiveness. Studies examining this association in the context of depression and HC have yielded equivocal findings (e.g. positive correlations in McRae et al. 2012a, no association in Dillon and Pizzagalli 2013; Wolgast et al. 2011). These findings cannot necessarily be generalized to psychosis because the absence of an association in depression might be driven by motivational aspects (e.g. principally being able to use reappraisal successfully, but not doing so habitually due to a lack of motivation). Even though motivational aspects might also be relevant in PD with negative symptoms, PD may also be more likely to experience difficulties in applying reappraisal successfully (e.g. due to neurocognitive deficits) and thus be less likely to use the strategy habitually. Whether there is such a correlation in PD therefore still needs to be examined.

Summing up, although PD report to use reappraisal to a lesser extent than HC, there is still a significant amount of uncertainty regarding potential deficits in the successful use of reappraisal. Also, it is unknown whether the habitual use of reappraisal is related to its success and
whether alexithymia, executive functions and a jumping-to-conclusion bias account for group differences regarding reappraisal success. Improving on several design aspects of previous research (e.g. sample size, inclusion of a clinical control group, focus on more than one emotion and assessment of subjective and psychophysiological indicators of emotion-regulation), we hypothesized that: (1a) PD report to use reappraisal less often compared to HC and patients with anxiety disorders (AD), and (1b) PD are less successful compared to HC and AD in reducing experimentally induced anxiety and sadness through reappraisal; (2) the self-reported habitual use of reappraisal is related to whether it can be applied successfully; (3a) alexithymia, (3b) neurocognitive deficits, and (3c) the jumping-to-conclusion bias mediate group differences in reappraisal success.

Methods

Procedure

The data was collected in a research project on emotion-regulation in psychosis (LI 1298/7-1) that was approved by the Ethical Committee of the German Psychological Society. Within this project, eligible participants (PD, individuals at clinical high risk for psychosis, AD and HC) were invited for a detailed assessment of inclusion criteria, sociodemographic information, clinical data, and habitual emotion-regulation skills.

On two additional assessment days, neuropsychological tests were conducted followed by a social exclusion paradigm and an emotion-regulation paradigm that were conducted in a randomized order. All participants were paid for participating in the study. Data related to the social exclusion paradigm (Lincoln et al. 2018; Sundag et al. 2018) and questionnaire data (Ascone et al., 2020) have been published elsewhere. Here, we analyze the data from the questionnaires and from the experimental paradigm on emotion-regulation. We use the data provided by the PD, AD, and HC and focus on reappraisal as the strategy of interest.

Participants

Participants were recruited from inpatient and outpatient treatment settings in and around Hamburg/Germany via online advertisement and advertisements in local newspapers/bulletin boards. Inclusion criteria for all participants were: (1) age between 18 and 65 years, (2) sufficient fluency in the German language, (3) ability to provide informed consent, (4) IQ > 85, (5) no acute suicidality, and (6) no neurological disorders, dementia, diagnosis of bipolar disorder or substance-dependence.

PD fulfilled the criteria for a diagnosis of a psychotic disorder according to the Mini International Neuropsychiatric Interview (M.I.N.I., Sheehan et al. 1998) either (a) acutely or within the previous two years, or (b) at any time of their life, if they had experienced at least two psychotic episodes, were currently taking antipsychotic medicine and scored either at least mildly on two items from the positive or negative subscale of the Positive and Negative Syndrome Scale (PANSS; score = 3; Kay, Fiszbein, & Opfer, 1987) or at least moderately on one item (score ≥ 4).

AD were required to fulfill criteria for at least one anxiety disorder (social phobia, generalized anxiety disorder or panic disorder with or without agoraphobia) in the M.I.N.I. at the time of assessment, but to not fulfill criteria for schizoaffective, delusional disorder.

HC were required to not fulfill criteria for any past or present axis-I disorder in the M.I.N.I., to not be taking medication for any type of mental problem and to not have a first-degree relative with a psychotic disorder.

Measures

Psychopathology

Symptom severity in PD was assessed with the Positive and Negative Syndrome Scale (PANSS; Kay et al. 1987). This semi-structured interview assesses positive symptoms (7 items), negative symptoms (7 items) and global psychopathology (16 items). Internal consistencies for this study are within an acceptable range (positive subscale: $\alpha = 0.67$; negative subscale: $\alpha = 0.70$; global psychopathology: $\alpha = 0.71$).

Habitual Reappraisal

The habitual use of reappraisal was measured with the positive reappraisal subscale of the Cognitive Emotion Regulation Questionnaire (CERQ; German version; Loch et al. 2011), a 36-item questionnaire designed to assess the habitual use of nine separate cognitive emotion-regulation strategies (positive reappraisal, acceptance, positive refocusing, refocus on planning, putting into perspective, self-blame, rumination, catastrophizing, and blaming others). It is rated on a 5-point scale ranging from 1 (almost never) to 5 (almost always). The internal consistency for the positive reappraisal subscale in our study is acceptable ($\alpha = 0.75$).

Alexithymia

Problems in identifying and describing one’s emotions were assessed with the German version of the Toronto Alexithymia Scale (TAS-20; Bach et al. 1996). It consists of 20 items that are rated on a 5-point scale ranging from 1 (not
Neurocognitive Functioning

The Trail-Making-Test (TMT; Reitan 1958) was conducted as an indicator of neurocognitive capacity (processing speed, visual search, scanning, cognitive flexibility, and executive functions). The TMT consists of two parts with TMT-A requiring participants to connect 25 encircled numbers scattered on a sheet of paper in ascending order and TMT-B requiring to connect numbers and letters in an alternative order (e.g. 1-A-2-B-...). The score is equivalent to the time in seconds needed to complete each task, respectively.

Jumping-to-Conclusion Bias (JTC)

JTC was assessed with the fish-task (Moritz et al. 2010), which is a revised and computerized version of the beads task (Huq et al. 1988). In this task, fish are caught one after another from one of two lakes that are filled with red and gray fish in opposing ratios (80% red and 20% grey). The participants have to decide from which of the two lakes the fish are caught and are allowed to request seeing as many fish as they like before deciding. There was no limit as to how many fish could be drawn. JTC was operationalized as making a decision after the first fish, as was done by Moritz and colleagues (2010).

Emotion-Regulation Paradigm

Reappraisal success was assessed with a commonly used experimental emotion-regulation paradigm in which negative affect is induced via affective stimuli and subsequently regulated via instructed emotion-regulation strategies. In this study, pictures from the International Affective Picture System (IAPS; Lang 2005) were used to induce anxiety and sadness (for normative ratings on discrete emotions and arousal see Appendix (Table 3)). During each trial, one of 16 anxiety-evoking and 16 sadness-evoking IAPS-pictures was presented for 9 s (induction phase, see Fig. 1). This was followed by an intensity-rating of the subjective emotion intensity (t1). Subsequently, an instruction was presented for 3 s informing the participants which emotion-regulation strategy to use (experimental condition: reappraisal, distraction, or acceptance) or instructing them to just view (control condition) while being presented with the same picture again for 12 s. This was followed by another intensity-rating of the subjective emotion intensity (t2).

To begin with, the participants were asked to sit still for a 4-min baseline heart rate (HR) measurement, and to rate their baseline anxiety and sadness level (t0). Then they completed a 10-min training phase in which instructions on how to apply the different emotion-regulation strategies where depicted on the screen followed by four anxiety eliciting pictures to practice each strategy and the control condition (just view). For reappraisal, the participants received the following instruction: “Look at the picture attentively. Try to change the emerging emotions by reappraising the situation”.

![Fig. 1 Structure of a trail within the experimental emotion-regulation paradigm (Color figure online)](image-url)
For the control condition (view), participants received the following instruction: “Look at the picture attentively. Concentrate on what is depicted on the picture. Do not apply another strategy (reappraisal, distraction, acceptance)” This was followed by the experimental paradigm, which was constructed as a randomized block design with each block consisting of four trials. The emotion-regulation paradigm was conducted as 4 × 2 × 4 mixed experimental design with emotion-regulation strategies (reappraisal, distraction, acceptance, view) and emotion (anxiety and sadness) as within subject factors and group as the between subject factor (PD, AD, individuals at clinical high risk for psychosis and HC). For the purpose of this study, we only used the reappraisal and the view condition.

The dependent variables were picture-congruent emotion intensity and physiological arousal. Emotion intensity was rated by the participant after each picture on well-validated 11-point rating scales (1 = “not applicable”, 11 = “applicable”; Stemmler et al. 2001). The scales were labeled with four adjectives selected to capture anxiety (frightened/timid/afraid/scared) and sadness (sad/depressed/miserable/dejected).

Physiological arousal, indicated by the skin conductance and HR, was recorded continuously during the experiment. The skin conductance was measured with a sampling rate of 32 Hz using the NeXus 32™ BioTrace™ system by Mind Media. Ag/AgCl electrodes were placed at the distal phalanx of the second and third finger of the left hand. The skin conductance data was processed using Matlab R2020a and Ledalab V3.4.9. Data was downsampled to 16 Hz and the minimum amplitude threshold was set to 0.01 μS. We used continuous decomposition analysis (Benedek and Kaernbach 2010) to extract the continuous phasic activity. For each trial we calculated the ‘area under the curve’ in terms of the integrated skin conductance response (ISCR; measured in μS*sec) as a representation of the “strength of affect” (Boucsein, 2012, p.165) separately for the emotion induction and the regulation phases. The time-frame of the analysis was 8 s, starting at 1 s after stimulus presentation, respectively, due to latency effects of the skin conductance response. ISCR-values were additionally log-transformed to obtain normal distribution. Participants were excluded from the analyses of skin conductance data in the case of insufficient quality/missing data and regular (= daily) substance abuse. The NeXus 32™ BioTrace™ system was also used to record HR. The blood volume pulse was measured from the index finger of the non-dominant hand with a sampling rate of 256/s. Visual artefact screening was performed. HR was averaged across 9 s in the induction phase and 12 s in the regulation phase. HR-values were additionally log-transformed to obtain normal distribution. Participants were excluded from the analyses of HR data in the case of insufficient quality/missing data, cardiovascular disease or anomalies or cardiac medication (e.g. beta-blockers, angiotensin-converting enzyme inhibitors). Participants stayed connected to the psychophysiological sensors throughout the entire emotion-regulation paradigm.

Statistical Analyses

Analyses were conducted in SPSS (version 24). To test for group differences in sample characteristics and habitual use of reappraisal, one-way analyses of variance (ANOVA) and χ²-tests were conducted, respectively. Where necessary due to a partial violation of assumptions, analyses were repeated with non-parametric tests (Kruskal–Wallis). To test for group differences in reappraisal success, separate mixed repeated measures ANOVAs were conducted for each dependent variable (subjective picture-congruent emotion intensity, ISCR and HR). Type of stimuli (anxiety vs. sadness inducing IAPS picture), timepoint (baseline, pre-regulation, post-regulation; for event-related ISCR only pre-regulation and post-regulation) and condition (reappraisal vs. view) were entered as within-subject factors and group (PD vs. AD vs. HC) was entered as the between-subject factor. The Greenhouse–Geisser adjustment was used to correct for violations of sphericity. Bonferroni adjustment was applied for post-hoc t-tests.

We expected the following effects: (1) a significant main effect for time with post-hoc tests showing a significant increase of picture-congruent emotion intensity and HR from baseline (t0) to post emotion induction (t1) across all groups, both conditions and both emotions (manipulation check); (2) a significant time*condition interaction with post-hoc tests showing no differences between conditions at baseline (t0) or after emotion induction (t1), but a significantly lower emotion intensity after reappraisal than after viewing (t2); (3) a significant time*condition*group interaction effect, indicating that the time*condition interaction effect described previously differs between groups. Specifically, we expected post-hoc tests to show a significantly higher emotion intensity after reappraisal in PD compared to HC and AD. In addition, we explored whether the type of emotion had an effect on potential reappraisal deficits in PD by also testing for a time*condition*group*emotion interaction.

In the initial analyses of the psychophysiological data (ISCR and HR), age, gender and cigarettes smoked per day were considered as covariates due to their potential effects on autonomic activity. Since no covariate reached significance, the final analyses were conducted without covariates.

Pearson correlation was used to test the interrelation between habitual use of reappraisal and reappraisal success (subjective success: difference in state anxiety and sadness from t1 to t2 within the reappraisal condition; physiological
indicators of reappraisal success: difference in ISCR and HR from t1 to t2 within the reappraisal condition).

Results

Sample Characteristics

Within the PD sample \((n=60)\), diagnoses included schizophrenia \((n=41)\), schizoaffective disorder \((n=15)\), delusional disorder \((n=3)\), and brief psychotic disorder \((n=1)\). The majority of PD \((n=57)\) received antipsychotic medication (chlorpromazine equivalence dose of \(M=437.60, SD=483.86\), calculated as proposed by Benkert and Hippius 2006). Within the AD sample \((n=40)\), disorders included social anxiety \((n=13)\), panic disorder with/without agoraphobia \((n=30)\), and generalized anxiety disorder \((n=10)\). Two AD reported to take antipsychotic medication (chlorpromazine equivalence dose of \(M=8.33, SD=11.79\)). PD did not differ from AD and HC regarding age, gender distribution and years of education. See Table 1 for means and statistics.

Group Differences in the Habitual Use of Reappraisal

The one-way ANOVA revealed significant group differences in habitual reappraisal, \(F(2, 137) = 8.085, p < 0.001; \eta^2_{\text{partial}} = 0.106\). Contrary to our hypothesis, pairwise-comparisons showed no significant difference in habitual reappraisal between PD and HC \((p=0.099)\). However, PD reported to use reappraisal significantly more often than AD \((p=0.008)\). Additionally, AD reported to use reappraisal significantly less often than HC \((p<0.001)\). See Table 2 for group-means and standard deviations.

Group Difference in Reappraisal Success

Results of reappraisal success for all outcome variables (emotion intensity, ISCR and HR) and both experimental conditions (reappraisal vs. view) are visualized in Figs. 2, 3 and 4. The following report is restricted to the effects that were hypothesized. An account of all effects is provided in the Appendix (Tables 4, 5, 6).

Table 1

| Variable                  | Group                      | Parameter tests | Non-parametric tests |
|---------------------------|----------------------------|-----------------|----------------------|
|                          | Patients with psychotic disorders \(n=60\) \((M, SD)\) | Patients with anxiety disorders \(n=40\) \((M, SD)\) | Healthy control subjects \(n=40\) \((M, SD)\) |                  |                  |
| Age in years              | 40.15 (11.66)              | 42.23 (11.64)   | 40.03 (10.78)        | \(F(2,137) = .497, p = .609\) | \(H(2) = 1.476, p = .478\) |
| Gender (%female)          | 63.3                       | 62.5            | 67.5                 | –                  | \(\chi^2(2) = .26, p = .878\) |
| Years of education        | 11.71 (1.57)               | 11.33 (1.72)    | 11.86 (1.53)         | \(F(2,137) = 1.224, p = .297\) | \(H(2) = 2.091, p = .352\) |
| PANSS positive            | 15.38 (5.28)               | –               | –                    |                    |                  |
| PANSS negative            | 11.98 (4.90)               | –               | –                    |                    |                  |
| PANSS general             | 30.45 (7.57)               | –               | –                    |                    |                  |

Table 2

| Variable               | Group                      | Parameter tests | Non-parametric tests |
|------------------------|----------------------------|-----------------|----------------------|
|                       | Patients with psychotic disorders \(M, SD)\) | Patients with anxiety disorders \(M, SD)\) | Healthy control subjects \(M, SD)\) | \(F(2, 137) = 8.085, p < .001\) | \(H(2) = 14.566, p = .001\) |
| CERQ reappraisal       | 8.83 (3.03)                | 7.23 (2.58)     | 9.83 (3.07)          | \(F(2, 137) = 25.031, p < .001\) | \(H(2) = 39.365, p < .001\) |
| TAS                    | 51.25 (10.35)              | 54.68 (11.15)   | 39.78 (8.01)         | \(F(2, 137) = 4.334, p = .015\) | \(H(2) = 8.426, p = .015\) |
| JTC bias (% yes)       | 41.7                       | 37.5            | 32.5                 | \(F(2, 137) = 8.259, p < .001\) | \(H(2) = 13.048, p = .001\) |
| TMT A                  | 31.21 (20.05)              | 27.25 (10.59)   | 22.43 (5.69)         | \(F(2, 137) = 4.334, p = .015\) | \(H(2) = 8.426, p = .015\) |
| TMT B                  | 61.32 (24.61)              | 54.22 (19.85)   | 44.16 (13.51)        | \(F(2, 137) = 8.259, p < .001\) | \(H(2) = 13.048, p = .001\) |

CERQ Cognitive Emotion Regulation Questionnaire (positive reappraisal subscale), TAS Toronto Alexithymia Scale, JTC jumping-to-conclusion bias measured with the fish-test, TMT Trail Making Test (version A and B)
Fig. 2 Graphical display of subjective emotion intensity within the reappraisal and control condition.

Fig. 3 Graphical display of integrated skin conductance response (ISCR; time integral of phasic driver over response window; log-transformed) within the reappraisal and control condition. After applying the additional exclusion criteria for the psychological data, the samples comprised \( n = 128 \) for the ISCR-analyses (PD: \( n = 53 \); AD: \( n = 39 \); HC: \( n = 36 \)).

Fig. 4 Graphical display of heart rate (HR; log-transformed) within the reappraisal and control condition. After applying the additional exclusion for the psychological data, the samples comprised \( n = 103 \) for the HR-analyses (PD: \( n = 44 \); AD: \( n = 28 \); HC: \( n = 31 \)).
Subjective Success

For the self-report of emotion intensity, the within-subject analysis revealed a significant main effect for time, $F(1.238, 169.562) = 42.993$, $p < 0.001$, $\eta^2_{\text{partial}} = 0.239$, with post hoc-tests showing a significant increase of picture-congruent emotion intensity from t0 to t1 across all groups, conditions and types of emotion elicited ($p < 0.001$), which indicates that emotion was successfully manipulated by the IAPS pictures. Negative emotions decreased from t1 to t2 ($p < 0.001$) but remained significantly higher at t2 compared to t0 ($p = 0.004$). A significant interaction effect was found for time*condition, $F(1.655, 226.667) = 77.973$, $p < 0.001$, $\eta^2_{\text{partial}} = 0.363$ with post-hoc tests indicating no differences in emotion intensity ratings between the conditions at t0 and t1 (both $p$s > 0.05), but significant differences between conditions at t2 with lower levels of emotion intensity in the reappraisal condition compared to the view condition ($p < 0.001$). Within the reappraisal condition, emotion intensity at t2 did not differ from t0 ($p > 0.999$). Within the viewing condition, emotion intensity at t2 was significantly higher compared to t0 ($p < 0.001$). No significant interaction effect was found for condition*time*group, $F(3.309, 226.667) = 1.229$, $p = 0.300$, $\eta^2_{\text{partial}} = 0.018$, indicating that PD showed a reduction of emotion intensity after reappraisal that was comparable to the reductions in both control groups. Finally, no significant interaction effect was found for condition*time*group*emotion, $F(3.701, 253.503) = 1.225$, $p = 0.301$, $\eta^2_{\text{partial}} = 0.018$.

Physiological Success

For ISCR as dependent variable, the within-subject analysis revealed a significant main effect for time, $F(1, 125) = 43.136$, $p < 0.001$, $\eta^2_{\text{partial}} = 0.268$, with post hoc-tests showing a significant decrease from t1 to t2 ($p < 0.001$). There was a significant interaction effect for time*condition, $F(1, 125) = 11.553$, $p = 0.001$, $\eta^2_{\text{partial}} = 0.085$, with post hoc-tests showing a higher ISCR within the view compared to the reappraisal condition at t1 ($p < 0.001$), but not at t2 ($p = 0.769$). ISCR reduced from t1 to t2 significantly in both the view condition ($p < 0.001$) and the reappraisal condition ($p = 0.026$). There was no significant interaction effect for condition*time*group, $F(2, 125) = 0.423$, $p = 0.656$, $\eta^2_{\text{partial}} = 0.007$, or for condition*time*group*emotion, $F(2, 125) = 0.124$, $p = 0.884$, $\eta^2_{\text{partial}} = 0.002$.

For HR as a dependent variable, the within-subject analysis revealed no significant main effect for time, $F(1.022, 102.244) = 0.271$, $p = 0.609$, $\eta^2_{\text{partial}} = 0.003$. There was no significant interaction effect for time*condition, $F(2, 200) = 0.296$, $p = 0.744$, $\eta^2_{\text{partial}} = 0.003$, for condition*time*group, $F(3.992, 199.577) = 1.092$, $p = 0.362$, $\eta^2_{\text{partial}} = 0.021$, or for condition*emotion*time*group, $F(3.818, 190.885) = 1.491$, $p = 0.209$, $\eta^2_{\text{partial}} = 0.029$.

Interrelation of Habitual Reappraisal Use and Reappraisal Success

The correlation between habitual reappraisal use and subjective reappraisal success across all groups and both types of emotion elicited was $r = 0.087$, $p = 0.309$ and remained non-significant when calculated per group (PD: $r = 0.108$, $p = 0.410$; AD: $r = 0.036$, $p = 0.828$; HC: $r = 0.111$, $p = 0.496$). Also, no significant correlations were found between the habitual reappraisal use and physiological indicators of reappraisal success, neither for ISCR (across all groups: $r = -0.012$, $p = 0.892$; PD: $r = 0.022$, $p = 0.872$; AD: $r = -0.247$, $p = 0.130$; HC: $r = -0.006$, $p = 0.972$) nor for HR (across all groups: $r = -0.122$, $p = 0.163$; PD: $r = -0.079$, $p = 0.558$; AD: $r = -0.117$, $p = 0.473$; HC: $r = -0.093$, $p = 0.589$).

Alexithymia, Cognitive Functioning and Jumping-to-Conclusion Bias as Mediators

As can be seen in Table 2, one-way ANOVAs revealed significant group differences for emotion awareness and TMT-A and TMT-B performance. Pairwise comparisons indicated that PD and AD reported more difficulties with emotion awareness than HC (both $p$s < 0.001), whereas PD and AD did not differ ($p = 0.286$). Pairwise comparisons indicated poorer TMT performance in PD compared to HC (TMT-A: $p = 0.012$; TMT-B: $p < 0.001$), but not compared to AD (TMT-A: $p = 0.564$; TMT-B: $p = 0.285$). No group differences were found for JTC.

The planned mediation analyses were discarded due to lack of group differences in reappraisal success and because there were no significant correlations between reappraisal success and alexithymia, cognitive functioning or JTC, respectively (all $p`s > 0.005$).

Discussion

The main focus of this study was to examine potential psychosis-specific deficits in the habitual use and the successful application of reappraisal.

Our hypothesis that PD would report to use less reappraisal than HC and AD was not supported by our data. Although our study is unlikely to fundamentally change the small to moderate effect in a recent meta-analysis that indicated less habitual reappraisal in PD (Ludwig et al. 2019), it adds to the existing heterogeneity. This raises the question of possible moderators. Differences in patient characteristics, such as the amount of psychological therapy or acuteness of
disorder might contribute to the differences found between studies and should be considered in further research. Interestingly, several recent experience sampling studies even indicate a higher use of reappraisal in PD compared to HC (Ludwig et al. 2020; Visser et al. 2018). We would argue that the findings from these studies are more reliable than the retrospective questionnaire assessments because they enable a timely assessment of the use of reappraisal in daily life and are less susceptible to recall biases. Thus, we may have been underestimating the habitual use of reappraisal in psychosis.

Regarding regulation success, our results indicate that PD are as successful as HC and AD in reducing subjective anxiety and sadness through reappraisal. Even though this was also unexpected, it corroborates several other studies finding no deficits in subjective reappraisal success in psychosis (Grezellschak et al. 2015; Painter et al. 2019; Perry et al. 2012). Since the symptomatology in our sample is comparable to other samples of PD seeking therapeutic treatment (e.g. Lincoln et al. 2012; Mehl et al. 2018), it is unlikely that the lack of deficits is explicable by having drawn a particularly healthy sample. Regarding the physiological indicators, we found a reduction of the skin conductance response in both the reappraisal and the control condition. The fact that these results do not mirror the results of the subjective ratings in all aspects is in line with other studies reporting discrepancies between self-reported and physiological arousal in psychosis (Söder et al. 2018). The failure to find an effect of reappraisal on HR and the skin conductance response is in line with a previous study (Westermann et al. 2014). PD showed a significantly higher HR than AD and HC across all timepoints [see Appendix (Table 6)], a pattern which has also been found in other studies (e.g. Bär et al. 2005; Lincoln et al. 2015; Rechl et al. 1994; Valkonen-Korhonen et al. 2003). Following emotional stimuli, an initial HR deceleration can be observed as an orienting response followed by a HR acceleration (Bradley 2009). Because we aimed to assess HR as a physiological indicator of ER success rather than to examine the emotional response itself, we examined the mean HR. This approach was also chosen in similar studies (e.g. Painter et al. 2019; Westermann et al. 2014). However, since HR did not increase in response to the emotion-eliciting images, it is difficult to interpret this measure as an indicator for reappraisal success.

Taken together, our study suggests that people with psychotic disorders can successfully use reappraisal, at least when they are instructed to do so in a laboratory setting. This does not rule out the possibility that the distress resulting from individually relevant and potentially psychosis-specific triggers in daily life might be more difficult to regulate. However, recent findings from an experience sampling study indicate intact reappraisal abilities in PD outside of the laboratory (Ludwig et al. 2020). Taken together, a growing number of studies do not support the notion that psychosis is associated with a deficit in using reappraisal successfully when operationalized in terms of subjective emotion experience. Nevertheless, studies using fMRI or EEG found differences between PD and HC in the modulation of neural responses (late positive potentials) through reappraisal (Horan et al. 2013; Strauss et al. 2013), and in the activation of brain regions associated with emotion-regulation (Modinos et al. 2010; Van Meer et al. 2014) that point towards regulation deficits within the neural circuits. The reasons for this discrepancy need to be further elucidated. Possibly, reappraisal deficits that are detectable on the neural level may be overridden by other processes (e.g. by compensatory mechanisms; Modinos et al. 2010; Phillips and Seidman 2008) and therefore do not translate into subjective experience, or a lack of emotion awareness contributes to the discrepancies between subjective report and physiological measures, as has been suggested in other studies (Söder et al. 2018). Another possibility concerns the time point when reappraisal is applied: two of the studies that found differences between PD and HC in the late positive potentials examined reappraisal as a antecedent-focused strategy by preceding emotion eliciting pictures with either negative or neutral descriptions (Horan et al. 2013; Strauss et al. 2013). In contrast, all studies focusing on the subjective success (including ours) asked participants to apply reappraisal during or after emotion generation, thus examining reappraisal as a response-focused strategy. Whether the timing of reappraisal might have an effect on regulation success has to be examined in future studies.

Surprisingly, self-reported use of habitual reappraisal and the ability to apply this strategy successfully were uncorrelated. As non-significance was found within each group, the missing association is unlikely to be due to disorder specific factors such as higher alexithymia in PD. Alternative explanations are that questionnaire-based estimates of habitual reappraisal are inaccurate, or that reappraisal success seen in a laboratory setting is not equivalent to reappraisal success in daily life. In any case, the missing association suggests that we cannot generalize findings of reduced habitual use of reappraisal from questionnaire-based studies to surmise general deficits in the use of that strategy. Rather, we may need to distinguish between different forms and levels on which deficits might occur, such as the frequency in the habitual use, the success in reducing negative affect on a subjective or a physiological level, or the ability to select an appropriate strategy depending on the context (see also McRae 2013).

Finally, our mediation hypotheses were also not supported because we did not find differences in reappraisal success between groups or correlations between reappraisal success and alexithymia, neurocognition and jumping-to-conclusion biases, respectively. However, a recent study suggests that neurocognition might be more relevant when subjects have
to flexibly choose emotion-regulation strategies than when they implement instructed strategies (Pruessner et al. 2020).

Limitations and Suggestions for Further Research

When interpreting the results on regulation success, it has to be considered that the subjective emotion intensity ratings might have been influenced either by social desirability or by expectancy effects, which is a general limitation of subjective measures. This is particularly likely because the participants were aware that they performed emotion regulation, which is difficult to mask in such experiments. Thus, those biases might have contributed to the reduction in subjectively reported negative affect we see in all groups following reappraisal. The possibility of such biases emphasizes the need to also consider physiological indicators that are less likely to be biased than subjective ratings. As we did not see group specific differences in the physiological indicators over time (no time*condition*group interactions), there is no reason to assume that social desirability or expectancy effects affected the ratings of PD specifically. Thus, they are unlikely to have had an impact on our main research question.

Even though all participants received specific instruction on which emotion-regulation strategy to use (or not to regulate), we do not know whether and to which extent those instructions were followed, and we cannot rule out that additional strategies were used. Future studies might benefit from either including specific questions to query whether instructions were followed or from instructing participants to think aloud while applying the strategies.

Conclusion and Therapeutic Implications

The prevalence of high levels of negative affect in PD points to some type of deficit in emotion-regulation in this patient population. However, contrary to the common assumption and despite prevalent problems with alexithymia and neurocognitive functioning, this study found no deficits in PD regarding the habitual and successful use of reappraisal after instruction. Nevertheless, it remains unclear, whether PD are able to use reappraisal successfully without instruction, whether they are able to select this strategy in appropriate situations and why most studies report that they use reappraisal less habitually. Those questions need to be addressed before disregarding the relevance of reappraisal in psychosis altogether. From a clinical perspective, the finding that PD are able to use reappraisal successfully after instruction and in controlled conditions is promising, because it suggests that they could be guided by therapists on how to use it successfully in everyday life.

Author Contributions TL and JS contributed to the study conception and design. JS performed material preparation and data collection. SO performed analyses and wrote the first draft of the manuscript. MR has contributed substantially to the analyses of the skin conductance data. All authors commented on previous versions of the manuscript. All authors read and approved the final manuscript.

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Compliance with Ethical Standards

Conflict of Interest Sandra Opoka, Johanna Sundag, Marcel Riehle and Tania Lincoln declare that they have no conflict of interest.

Ethical Approval All procedures were in accordance with the ethical standards of the Ethical Committee of the German Psychological Society and with the 1964 Helsinki declaration and its later amendments.

Informed Consent All participants provided written informed consent.

Animal Rights No animal studies were carried out by the authors for this article.

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Appendices

Appendix A

See Table 3
Table 3  Selected IAPS pictures with ratings for sadness, anxiety and arousal

| IAPS No | Sadness M | SD | Anxiety M | SD | Arousal M | SD |
|---------|-----------|----|-----------|----|-----------|----|
| Anxiety |           |    |           |    |           |    |
| 1022    | 1.96      | 1.61| 5.31      | 2.74| 3.73      | 2.38|
| 1052    | 1.73      | 1.30| 6.18      | 3.21| 4.88      | 2.63|
| 1301    | 2.96      | 2.36| 5.30      | 2.91| 3.96      | 2.72|
| 1930    | 2.81      | 2.30| 6.04      | 2.63| 4.85      | 2.62|
| 1931    | 2.16      | 1.67| 5.72      | 2.29| 5.22      | 2.50|
| 5972    | 5.33      | 2.53| 6.37      | 2.45| 4.56      | 2.94|
| 1120    | 2.41      | 1.89| 6.41      | 2.66| 4.30      | 2.73|
| 1019    | 3.28      | 2.41| 3.13      | 2.48| 3.53      | 2.29|
| 1050    | 2.15      | 1.96| 6.00      | 2.78| 4.35      | 2.54|
| 1051    | 2.23      | 1.90| 5.23      | 2.83| 3.88      | 2.41|
| 1120    | 2.14      | 1.89| 6.41      | 2.66| 4.30      | 2.73|
| 1200    | 1.08      | 0.28| 5.60      | 2.69| 3.24      | 2.50|
| 1201    | 2.08      | 1.99| 6.39      | 2.73| 4.76      | 2.90|
| 1220    | 1.73      | 1.25| 5.27      | 2.75| 3.68      | 2.50|
| 1300    | 2.52      | 1.71| 5.96      | 2.42| 3.80      | 2.31|
| 6370    | 3.81      | 2.58| 6.65      | 2.42| 4.32      | 2.90|
| Sadness |           |    |           |    |           |    |
| 2053    | 5.92      | 2.72| 3.25      | 2.56| 5.26      | 2.47|
| 2141    | 7.00      | 2.47| 4.16      | 2.67| 3.16      | 2.17|
| 2205    | 7.38      | 1.84| 3.41      | 2.63| 3.67      | 2.46|
| 2276    | 6.60      | 2.27| 2.32      | 1.86| 3.00      | 2.16|
| 2800    | 7.63      | 1.66| 2.96      | 2.42| 5.79      | 2.54|
| 2900    | 6.35      | 2.35| 2.81      | 2.61| 3.62      | 2.29|
| 3220    | 6.71      | 2.08| 3.50      | 2.55| 4.44      | 2.45|
| 3230    | 6.97      | 2.37| 4.08      | 2.53| 3.90      | 2.58|
| 3300    | 6.29      | 2.39| 2.58      | 2.26| 3.95      | 2.43|
| 3301    | 6.85      | 2.36| 3.85      | 2.66| 3.61      | 2.65|
| 6570    | 6.23      | 2.55| 3.88      | 2.85| 5.21      | 2.94|
| 9041    | 6.08      | 2.16| 2.44      | 2.06| 2.84      | 1.99|
| 9252    | 6.08      | 2.12| 3.80      | 3.01| 3.56      | 2.40|
| 9421    | 7.16      | 2.01| 6.04      | 2.77| 3.60      | 2.65|
| 9561    | 7.49      | 1.78| 2.04      | 2.01| 3.84      | 2.45|
| 9921    | 6.82      | 2.49| 5.62      | 2.56| 4.80      | 2.85|

Selected from https://psychonomic.org/archive. Ratings from Libkuman et al. (2007)
### Appendix B

See Table 4

| Effects                  | df | F     | p   | η²partial |
|--------------------------|----|-------|-----|-----------|
| Main effects             |    |       |     |           |
| Condition                | 1.000 | 36.499 | <.001 | 0.210     |
| Emotion                  | 1.000 | 21.049 | <.001 | 0.133     |
| Time                     | 1.238 | 42.992 | <.001 | 0.239     |
| Group                    | 2    | 2.893 | .059 | 0.041     |
| Interaction effects      |    |       |     |           |
| Condition*group          | 2.000 | 0.304 | .738 | 0.004     |
| Emotion*group            | 2.000 | 1.012 | .366 | 0.015     |
| Time*group               | 2.475 | 3.121 | .036 | 0.044     |
| Condition*emotion        | 1.000 | 0.401 | .528 | 0.003     |
| Condition*emotion*group  | 2.000 | 0.423 | .656 | 0.006     |
| Condition*time           | 1.655 | 77.973 | <.001 | 0.363     |
| Condition*time*group     | 3.309 | 1.229 | .300 | 0.018     |
| Emotion*time             | 1.221 | 22.079 | <.001 | 0.139     |
| Emotion*time*group       | 2.443 | 1.568 | .206 | 0.022     |
| Condition*emotion*time   | 1.850 | 1.422 | .243 | 0.010     |
| Condition*emotion*time*group | 3.701 | 1.225 | .301 | 0.018     |

### Appendix C

See Table 5

| Effects                  | df | F     | p   | η²partial |
|--------------------------|----|-------|-----|-----------|
| Main effects             |    |       |     |           |
| Condition                | 1    | 6.361 | .013 | 0.048     |
| Emotion                  | 1    | 2.610 | .109 | 0.020     |
| Time                     | 1    | 43.136 | <.001 | 0.257     |
| Group                    | 2    | 0.485 | .617 | 0.008     |
| Interaction effects      |    |       |     |           |
| Condition*group          | 2    | 0.219 | .804 | 0.003     |
| Emotion*group            | 2    | 1.813 | .167 | 0.028     |
| Time*group               | 2    | 0.289 | .750 | 0.005     |
| Condition*emotion        | 1    | 5.693 | .019 | 0.044     |
| Condition*emotion*group  | 2    | 0.043 | .958 | 0.001     |
| Condition*time           | 1    | 11.553 | .001 | 0.085     |
| Condition*time*group     | 2    | 0.423 | .656 | 0.007     |
| Emotion*time             | 1    | 1.290 | .258 | 0.010     |
| Emotion*time*group       | 2    | 0.257 | .774 | 0.004     |
| Condition*emotion*time   | 1    | 6.152 | .014 | 0.047     |
| Condition*emotion*time*group | 2    | 0.124 | .884 | 0.002     |

After applying the additional exclusion criteria for the skin conductance analyses, the sample comprised n = 128 (PD: n = 53, AD: n = 39, HC: n = 36)

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