Review article

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Clinical implications of fear extinction in anxiety disorders

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Abstract: Anxiety disorders (ADs) are characterized by increased chronicity and comorbidity with other ADs. Although exposure is the most effective therapy option for ADs, some patients show poor treatment response and a heightened vulnerability for relapse after treatment completion. Hence, significant research effort needs to be devoted to improve the long-term effectiveness of exposure effects. Recent attempts to increase exposure therapy efficacy use strategies aimed at promoting the acquisition and retrieval of extinction memories. The present review illustrates the value and limitations of such extinction-based therapy approaches. We present and discuss recent findings from translational studies using cortisol and self-efficacy enhancement as an add-on to exposure therapy. We illustrate how the integration of findings from experimental research on fear extinction learning and self-efficacy could advance the development of more optimized treatments for ADs.

Keywords: cortisol; exposure therapy; fear extinction; self-efficacy; therapy generalization.


Schlüsselwörter: Furchtextinktion; Generalisierungseffekte; Cortisol; Selbstwirksamkeit; Konfrontationstherapie.

Anxiety disorders (ADs) belong to the most prevalent mental disorders (Bandelow and Michaelis, 2015). Chronicity and comorbidities with other ADs affect the disease course of ADs (Bandelow and Michaelis, 2015). Cognitive behavioral therapy (CBT) is both highly efficient and effective in the treatment of ADs (Otte, 2011). CBT involves a set of cognitive and behavioral interventions such as exposure. Although exposure is the most effective therapeutic tool for ADs, some patients fail to exhibit significant symptom improvement or show recovery of fear and avoidance after completion of exposure (Hoffmann and Smits, 2008; Norton and Price, 2007). Current knowledge on the mechanisms governing the beneficial effects of exposure has been largely influenced by the general propositions of the inhibitory learning and inhibitory regulation models (Craske et al., 2006, 2008). Here, fear extinction is considered a central candidate to explain the beneficial effects of exposure as well as relapse phenomena after successful treatment (Craske and Mystkowski, 2006).
The association between fear extinction and exposure therapy

From an inhibitory learning perspective (Craske and Mystkowski, 2006, 2008), interindividual variability in exposure treatment outcome (Norton and Price, 2007) may be explained by the level of interindividual differences in fear extinction learning. In line with this idea, initial studies demonstrated that differences in extinction learning performance during a differential fear conditioning task were associated with variability in CBT outcomes in phobic children (Waters and Pine, 2016) and with the level of treatment gains during an exposure therapy analog in spider phobia (Forcadell et al., 2017). Likewise, anxiety reductions after exposure in patients with social anxiety could be predicted on the basis of extinction learning performance (Ball et al., 2017). Work from our group indicates that interindividual differences in fear extinction learning might also influence the patient’s general ability to complete an exposure task within one or two sessions (Raeder et al., 2020). We showed that spider-phobic participants who were able to complete exposure within two 60-min sessions (i.e., completers) exhibited more pronounced short- and long-term therapy benefit than non-completers. Most importantly, fear extinction performance is linked to the ability to complete the exposure. Completers showed more pronounced fear extinction (retrieval) relative to noncompleters. This finding indicates that one subgroup of patients with specific phobia (non-completers) failed to accomplish exposure in a predetermined time possibly owing to deficient fear extinction.

These results bear important implications for the implementation of exposure in routine care. The inability of some patients to accomplish exposure in a predetermined time might be at odds with the specific regularities of routine care (Gunther and Whittal, 2010). To conclude, research on fear extinction might not only explain the variability in exposure treatment efficacy across patients but also bear specific implications for the implementation of exposure to routine care (Richter et al., 2017).

Pharmacological enhancement of exposure therapy efficacy: lessons from studies using cortisol as an add-on to exposure therapy

Studying exposure treatment processes from the perspective of the fear extinction model might provide valuable information on how to optimize exposure treatment efficacy (Craske et al., 2018). A great number of studies showed alterations in fear acquisition (Mosig et al., 2014) and/or deficits in fear extinction in ADs (Lissek et al., 2005). Fortunately, research on cognitive and neurobiological mechanisms of extinction produced a great wealth of meaningful results on how fear extinction can be selectively enhanced (Craske et al., 2018). For instance, data from animal and human work suggest that stress and cortisol can modulate the acquisition, consolidation, and retrieval of extinction memories (de Quervain et al., 2017; Stockhorst and Antov, 2016). Accordingly, the translation of these findings to the context of exposure therapy has received great interest. Systemic administration of glucocorticoids (i.e., cortisol) before exposure has been shown to enhance the efficacy of exposure-based treatments (for a review, see de Quervain et al., 2017). However, possible timing-dependent effects of cortisol on exposure outcome have been neglected in clinical studies. This is surprising given that the effects of stress and cortisol on (extinction) memory processes are not ubiquitous but might depend on the exact timing of administration (Stockhorst and Antov, 2016). Likewise, existing clinical studies rarely considered the possible impact of cortisol on the generalization of therapy effects across contexts. Given the existence of context-specific effects of cortisol on fear extinction (Meir Drexler et al., 2019), cortisol might affect the generalization of exposure treatment effects from the treatment context to other unfamiliar contexts. In other words, owing to the impact of cortisol on context specificity of fear extinction, the pharmacological enhancement of exposure with cortisol can possibly aggravate or dampen the return of fear after successful treatment. In the therapy setting, return of fear can be observed when patients encounter their feared object in an unfamiliar context, for example, seeing a spider in the basement instead of the room in which the exposure therapy took place – termed as fear renewal (Craske and Mystkowski, 2006, 2008).

Raeder et al. (2019a) conducted the first study that examined the effect of cortisol on both exposure efficacy and fear renewal after exposure. We showed that the administration of cortisol after exposure did not enhance the efficacy of exposure therapy in spider phobia. We further observed a detrimental effect on context-dependent return of fear (fear renewal) in the long term in participants who received cortisol relative to placebo-treated participants. In light of previous clinical studies in this field (de Quervain et al., 2017), our findings indicate that cortisol may boost exposure therapy efficacy only when given before (rather than after) exposure. Furthermore, the exact timing of cortisol administration seems to be critical when attempting to increase the generalization of therapeutic effects across contexts (Meir Drexler et al., 2019). Accordingly, post-exposure
administration of cortisol might be a less well-suited augmentation strategy because it may lead to an increase in fear renewal in the long term.

**Exposure treatment efficacy: mediators and moderators**

Some important mediators/moderators need to be taken into consideration when conducting clinical studies assessing the putative role of cognitive enhancers in exposure therapy. ADs are more frequent in women (Kessler et al., 2005). There is a gender-dependent effect in fear extinction (Merz et al., 2018). The sex hormone estrogen may affect the short- and long-term processing of extinction memories (Maeng and Milad, 2015). The use of oral contraceptives (OCs) affects endogenous estrogen secretion in women. Women using OCs show impaired fear extinction learning (Merz et al., 2012, 2018). Considering the influence of OC use and the variations in estrogen levels during the menstrual cycle in the context of exposure treatment in women might therefore be highly valuable. In support of this proposition, work from our laboratory (Raeder et al., 2019c) and that from others (Graham et al., 2018) indicate that hormonal contraceptive use in women has an impact on the immediate and long-term effects of exposure. Precisely, free-cycling women and women using hormonal contraceptives showed different response profiles to exposure therapy (Raeder et al., 2019c). Spider-phobic women using hormonal contraceptives exhibited less fear reduction and symptom improvement from pre-treatment to post-treatment and at six-week follow-up than their free-cycling counterparts.

The aforementioned findings suggest that the implementation of hormonal measurements and the systematic assessment of contraceptive use, which itself affects variability in exposure outcome, is important to derive a complete picture on the possible effects of cognitive enhancers in exposure therapy. This is especially true with regard to the effects of cortisol because OC use alters the effects of cortisol on fear learning (Merz et al., 2012, 2018). Interestingly, the Stress Timing affects Relapse (STAR) model has been proposed as a valuable framework to stimulate future clinical studies on the interaction between cortisol and sex hormones on extinction memories (Meir Drexler et al., 2019).

**Exposure: is it more than fear extinction?**

The rationale behind exposure therapy is to assist patients in overcoming their anxiety by creating a safe environment in which they encounter feared or avoided scenarios. Thus, a central goal of exposure is to induce positive mastery experiences that are ideally accompanied by substantial decrements of fear and avoidance in treated patients. According to Bandura (1988), positive mastery experiences lead to an increase in self-efficacy beliefs, which might constitute a prerequisite for a successful CBT. Several studies showed a positive association between increased self-efficacy and therapy outcome in patients with ADs (Bouchard et al., 2007; Gallagher et al., 2013). Given the mutual relationship between exposure and self-efficacy, the selective modulation of perceived self-efficacy might be effective to promote key processes (i.e., fear extinction learning) of exposure. In line with this, we showed that an increase in perceived self-efficacy (induced by false-positive verbal feedback) affects the acquisition and retrieval of extinction memories. Healthy participants with an increased self-efficacy showed better fear extinction learning and retrieval in a differential fear conditioning task (Zlomuzica et al., 2015).

Promoting self-efficacy might also represent an effective strategy to increase exposure efficacy. Raeder et al. (2019b) recently showed that increasing self-efficacy via the active rehearsal of personal mastery experiences is suitable to promote exposure outcome in patients with height phobia. In particular, self-efficacy enhancement led to more pronounced reductions in fear and avoidance after one session of standardized exposure in virtual reality (Raeder et al., 2019b). The mechanisms underlying the beneficial effect of an increased self-efficacy on fear extinction remain to be explored. Increased self-efficacy might lead to changes in the processing of extinguished memories (Zlomuzica et al., 2015). Alternatively, the utilization of positive personal experiences might affect the way how subjects perceive and cope with future challenges (Margraf and Zlomuzica, 2015; Zlomuzica et al., 2018). Since an adaptive processing of mastery experiences is fundamental to self-efficacy (Bandura, 1988), a better understanding of mechanisms underlying the storage and retrieval of personally relevant memories in patients with ADs would be highly valuable (Zlomuzica et al., 2014, 2016).

**Generalization of exposure therapy effects**

The comorbidity of ADs with other ADs is common (Bandelow and Michaelis, 2015). For instance, phobic individuals tend to suffer from multiple fears at the same time
Individuals with fear of spiders tend to fear other similar insects (e.g., cockroaches) and/or small animals (e.g., rats). Such multiple fears might even be functionally related and lead to an increase in psychopathological symptoms (Rachman and Lopatka, 1986a, b). Surprisingly, recommendations on how to systematically treat multiple fears and/or comorbid anxieties do not exist. Likewise, there is no therapeutic tool that can induce a generalization of therapeutic effects for different functionally related fears. Notwithstanding, we have recently shown that exposure might lead to a generalization of therapeutic effects to untreated fear stimuli. In particular, Preusser et al. (2017) demonstrated that exposure-induced reduction in fear and avoidance can also be observed for untreated stimuli, that is, those that bear feature overlap with treated stimuli but do not belong to the same category of fear stimuli (Figure 1).

These findings, which present to our knowledge the first study on this research gap, indicate that exposure effects are not restricted to the specific fear stimulus used during exposure (Preusser et al., 2017). Interestingly, such a generalization of clinical exposure treatment was recently also demonstrated in other fears (Hollander et al., 2020). How can we explain such generalization of exposure therapy effects? Findings from basic research on fear generalization (in particular, extinction generalization) (Dymond et al. 2015; Pittig et al., 2018) cannot fully account for the generalization of exposure effects to untreated fear stimuli. Alternatively, the self-efficacy concept of Bandura offers a more parsimonious account for the generalization of mastery experiences across different related (fear) domains (Bandura, 1988). Nevertheless, studying the generalization of therapeutic effects in ADs represents an important but neglected research field (Pittig et al., 2018).

Conclusions

Fear extinction might be a central candidate to explain exposure therapy benefit. The formation of personal mastery experiences during exposure leads to an increased self-efficacy, which might constitute another important element of a successful therapy for ADs. Attempts to promote fear extinction learning (e.g., via pharmacological modulation with cognitive enhancers) and to increase self-efficacy represent promising strategies to enhance exposure treatment efficacy and increase generalization of therapeutic effects.

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