Pharmacology of cognitive enhancers for exposure-based therapy of fear, anxiety and trauma-related disorders

N. Singewald a,⁎, C. Schmuckermair a, C. Whittle a, A. Holmes b, K.J. Ressler c

a Department of Pharmacology and Toxicology, Institute of Pharmacy and CMBI, Leopold-Franzens University of Innsbruck, Innrain 80-82, A-6020 Innsbruck, Austria
b Laboratory of Behavioral and Genomic Neuroscience, National Institute on Alcohol Abuse and Alcoholism, NIH, Bethesda, MD, USA
c Department of Psychiatry and Behavioral Sciences, Emory University School of Medicine, Atlanta, GA, USA

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A B S T R A C T
Pathological fear and anxiety are highly debilitating and, despite considerable advances in psychotherapy and pharmacotherapy, they remain insufficiently treated in many patients with PTSD, phobias, panic and other anxiety disorders. Increasing preclinical and clinical evidence indicates that pharmacological treatments including cognitive enhancers, when given as adjuncts to psychotherapeutic approaches [cognitive behavioral therapy including extinction-based exposure therapy] enhance treatment efficacy, while using anxiolytics such as benzodiazepines as adjuncts can undermine long-term treatment success. The purpose of this review is to outline the literature showing how pharmacological interventions targeting neurotransmitter systems including serotonin, dopamine, noradrenaline, histamine, glutamate, GABA, cannabinoids, neuropeptides (oxytocin, neuropeptides Y and S, opioids) and other targets (neurotrophins BDNF and FGF2, gluco corticoids, L-type-calcium channels, epigenetic modifications) as well as their downstream signaling pathways, can augment fear extinction and strengthen extinction memory persistently in preclinical models. Particularly promising approaches are discussed in regard to their effects on specific aspects of fear extinction namely, acquisition, consolidation and retrieval, including long-term protection from return of fear (relapse) phenomena like spontaneous recovery, reinstatement and renewal of fear. We also highlight the promising translational value of the preclinical research and the clinical potential of targeting certain neurochemical systems with, for example α-cyclodextrin, yohimbine, cortisol, and L-DOPA. The current body of research reveals important new insights into the neurobiology and neurochemistry of fear extinction and holds significant promise for pharmacologically-augmented psychotherapy as an improved approach to treat trauma and anxiety-related disorders in a more efficient and persistent way promoting enhanced symptom remission and recovery.

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Abbreviations: 5-HT, 5-hydroxytryptamine = serotonin; AC, adenylate cyclase; AMPA, α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid; AMY, amygdala; BA, basal amygdala; BDNF, brain-derived neurotrophic factor; BLC, basolateral complex; BLA, basolateral amygdala; CaMKII, Ca2+/calmodulin-dependent protein kinase II; CREB, cAMP response element binding; CNS, central nervous system; CREB, cAMP response element binding; CS, conditioned stimulus; DA, dopamine; D-cycloserine, yohimbine, cortisol, and L-DOPA. The current body of research reveals important new insights into the neurobiology and neurochemistry of fear extinction and holds significant promise for pharmacologically-augmented psychotherapy as an improved approach to treat trauma and anxiety-related disorders in a more efficient and persistent way promoting enhanced symptom remission and recovery.

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1. Introduction

Fear, anxiety and trauma-related disorders are associated with excessive fear reactions triggered by specific objects, situations or internal and external cues in the absence of any actual danger, and often include an inability to extinguish learned fear and to show adequate safety learning ([Jovanovic et al., 2012; Michael et al., 2007; Milad et al., 2009; Milad et al., 2013; Wessa & Flor, 2007] reviewed in [Homes & Singewald, 2013]) and ([Kong et al., 2014]). Pathological fear and anxiety occur in a range of psychiatric conditions, including various types of phobia (e.g. social phobia, agoraphobia or specific phobia), panic disorder with/without agoraphobia, obsessive-compulsive disorder (OCD), generalized anxiety (GAD) and post-traumatic stress disorder (PTSD) (DSM-5, 2013; ICD-10, 1994). These disorders comprise the most common mental disorders and are estimated to have a life-time prevalence of up to 28% among western populations (Kessler et al., 2005; Kessler et al., 2012; Wittchen et al., 2011). In addition to the personal suffering of patients, the economic burden caused by anxiety disorders is heavy (Gustavsson et al., 2011).

Available pharmacological and psychotherapeutic treatments (Bandelow et al., 2007) which aim to reduce fear and anxiety are associated with decreased symptom severity, but up to 40% of anxiety patients show only partial long-term benefit, and a majority of them fail to achieve complete remission (Bandelow et al., 2012; Hoffman & Mathew, 2008; Stein et al., 2009) clearly underlining the need for further improvement. Current pharmacological approaches either induce rapid anxiolytic effects (e.g. benzodiazepines, some antipsychotics) or require prolonged, chronic treatment (e.g. antidepressants) to attenuate symptoms of pathological fear and anxiety. Commonly employed psychotherapeutic interventions apply cognitive behavioral strategies and exposure techniques to help patients overcome the maladaptive beliefs and avoidance behaviors that reinforce the pathology related to fear-eliciting cues. Meta-analyses show that cognitive behavioral therapy (CBT) does have efficacy for several anxiety disorders, including PTSD, but patients have difficulty bearing the demanding and exhausting process of therapy and many who do manage to cope with it respond only partially and often relapse with time (Choy et al., 2007).

One strategy to improve CBT is to augment psychotherapy with adjunctive pharmacological treatments. Early attempts at combining ‘CBT’ with anxiolytic medications (e.g. benzodiazepines (BZD)) showed that the combination was no more effective [in some instances even counterproductive (Marks et al., 1993; Wilhelm & Roth, 1997)] than psycho- or pharmacotherapy alone [for details see (Dunlop et al., 2012; Hofmann, 2012; Otto et al., 2010a,b; Rodrigo et al., 2011)]. However, at least in some cases, this failure may have reflected idiosyncratic effects of the drugs tested (especially BZDs) rather than utility of the strategy itself, and there has been an intense search to identify agents that serve as more effective adjuncts to CBT. The preclinical assay most frequently used in this search is fear extinction — the focus of this current review. Extinction of fear following Pavlovian fear learning (Pavlov, 1927) in animals is procedurally similar to exposure-based CBT (Milad & Quirk, 2012). We will briefly outline different aspects of Pavlovian fear learning [(which is thought to be involved in the etiology and maintenance of anxiety disorders, e.g. (Amstadter et al., 2009)] and extinction highlighting the key processes that could be targeted to augment fear extinction (see Fig. 1 for an overview).

1.1. Fear and fear extinction

Experimentally, fear conditioning occurs when a previously neutral stimulus (conditioned stimulus (CS) — such as a tone or light) is paired with an aversive, unconditioned stimulus (US — e.g. electric shock to the forearm in humans, mild foot shock in rodents), resulting in a CS–US association whereby the CS alone elicits a conditioned fear response (e.g. freezing in rodents or increased skin conductance in humans). Following a successful CS–US association, fear memories require consolidation, a process involving a cascade of molecular and cellular events that alter synaptic efficacy, as well as a prolonged systems level interaction between brain regions, to stabilize the memory (McGaugh, 2000). Once consolidated, fear memories, reactivated by presenting the CS, are destabilized to render the original fear memory liable to pharmacological/behavioral interference (see overview in Fig. 1) and this is then followed by a second phase of molecular and cellular events to re-stabilize (re-consolidate) the (adapted) memory (Nader et al., 2000). Fear memories can be attenuated by various processes and interventions (including pharmacological and psychological approaches), some producing temporary blunting of fear behaviors and others causing more long-lasting relief. Interfering with the re-consolidation by inhibiting molecular and cellular events supporting fear memory re-stabilization [Fig. 1C, (Lee et al., 2006; Nader et al., 2000)], for example with β-adrenoceptor blockers (Debiec & Ledoux, 2004; Kindt et al., 2009), has been proposed as a clinical approach to alleviating fear memories. For discussion on this and other means of reducing fear (e.g. via US habituation) we refer the reader to some excellent prior reviews (Graham et al., 2011; Schwabe et al., 2014). Other potential ways to relieve fear include safety learning (Kong et al., 2014; Rogan et al., 2005) and erasure-like mechanisms such as destruction of erasure-preventing perineuronal nets (Gogolla et al., 2009).

Alternatively, fear memories can also be extinguished. Fear extinction, a process originally described by Pavlov (Pavlov, 1927), entails repeated exposure to anxiety-provoking cues to establish a new memory that counters the original fear memory. The process is highly relevant to fear, anxiety and trauma-related disorders which are associated with negative emotional reactions triggered by specific objects, situations or internal and external cues that are excessive to the actual danger posed. Moreover, extinction in animals is procedurally similar to forms of CBT that rely on exposure to anxiety-provoking cues (see Fig. 1) (Milad & Quirk, 2012), and anxiety disorders are associated with an inability to extinguish learned fear and to respond adequately to safety signals (Jovanovic et al., 2012; Michael et al., 2007; Milad et al., 2009; Milad et al., 2013; Wessa & Flor, 2007) [reviewed in (Homes & Singewald, 2013)]. Thus, fear extinction has considerable translational utility. The key processes that can be targeted to pharmacologically augment fear extinction are summarized in Fig. 1.

Extinction is a learning process driven by violation of the original CS=US contingency (termed ‘prediction error’ for review see (Pearce...
that extinction memories are prone to re-emergence (due to insufficient 'longterm extinction') indicates that the original fear memory is still in place and the extinction memory is weaker/more labile than the fear memory. This may be particularly true of older ('remote') fear memories (Tsai & Graff, 2014). The re-emergence of extinguished fear occurs under multiple circumstances: (i) renewal, when the CS is presented in a different context to that in which extinction training occurred; (ii) reinstatement, when the original US or another stressor is given unexpectedly; and (iii) spontaneous recovery, when a significant period of time has elapsed following successful extinction training (Henry et al., 2010; Myers & Davis, 2007). The likelihood of fear re-emergence is dependent on the strength of the extinction memory which is also determined by the type of extinction protocol used see e.g. (Laborda & Miller, 2013; Li & Westbrook, 2008). Spontaneous recovery (Rowe & Craske, 1998a,b; Schiller et al., 2008), reinstatement (Schiller et al., 2008) and renewal (Efting & Kindt, 2007) are all observable in clinical settings, and can be readily exploited in the laboratory to identify drugs and other interventions that can prevent fear re-emergence in animals and relapse in humans (Vervliet et al., 2013).

& Bouton, 2001; McNally & Westbrook, 2006), but it also contains other elements including habituation and desensitization some also say erasure/destabilization (Lin et al., 2011). Some authors therefore favor the term 'relearning' over extinction [for discussion see (Riebe et al., 2012)]. As with fear learning, extinction occurs in two phases: extinction acquisition and extinction consolidation. The decrement in the fear response during extinction training is also termed 'within-session extinction’. Similar to fear memory, stabilization of the extinction memory requires both a cascade of overlapping, but dissociable, molecular and cellular events [for review see (Myers & Davis, 2007; Orsini & Maren, 2012)] that alter synaptic efficacy and brain systems-level interactions (Pape & Pare, 2010). The strength of extinction memory can be assessed at some interval (usually >1 day) after extinction training in ‘extinction retrieval’ sessions (also termed ‘extinction retention’ or ‘extinction expression’, but note that we use the term ‘extinction retrieval’ throughout this review).

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Box 1
Why pharmacological augmentation of extinction?

A limitation of extinction-based exposure therapy is that patients can often relapse with the passage of time, with changes in context (out of the therapy context), or under conditions of stress or other provocations, such as experiencing trauma reminders. In the parable of learning theory, extinction memories are labile and fragile. A key goal for pharmacotherapy, therefore, is to identify compounds that overcome this fragility, by bolstering the formation, persistence and possibly context independence of extinction memories. One approach to achieving this goal is to use drugs as adjuncts to exposure therapy (cognitive enhancers) as a way to augment the extinction learning process. In this review, we discuss the various neurochemical and molecular signaling pathways that have been targeted to this end. On the one hand, there remain significant challenges to overcome, including the selective targeting of extinction-related processes without concurrent effects on original fear memories. On the other hand, there are clearly a plethora of potentially promising avenues to pursue, and we are optimistic that real advances can be made in treating trauma-related disorders.

Our goal in the current review is to offer a comprehensive overview of preclinical work on the possible pharmacological approaches (see Box 1 for an overview) to augment fear extinction and protect against the re-emergence of fear, and to discuss the potential translational value of these candidates as adjuncts to exposure-based CBT in anxiety patients.

2. Neuronal substrates of fear extinction

Using a number of complementary techniques (including electrophysiology, immediate-early gene mapping, tracing studies, lesioning/ inactivation approaches and optogenetics) research is revealing the complex and interconnected brain circuitry mediating fear extinction. Several key brain areas including the amygdala (AMY), hippocampus (HPC), medial prefrontal cortex (mPFC), periaqueductal gray (PAG), bed nucleus of the stria terminalis and others have been implicated in extinction [for recent detailed reviews see (Duvarci & Pare, 2014; Ehrlich et al., 2009; Herry et al., 2010; Knapska et al., 2012; Myers & Davis, 2007; Orsini & Maren, 2012; Pape & Pare, 2010)]. Of these, we focus on the AMY, HPC and mPFC as major, well-defined components of the fear circuitry (see Fig. 2).

While the AMY and the mPFC are crucial for the formation and maintenance of fear extinction memories, the HPC, linked with the mPFC and the AMY (Pape & Pare, 2010), processes contextual information linked with extinction (Orsini & Maren, 2012). Altering these hippocampal contributions to extinction memory or mimicking the hippocampal response within the extinction context (see below) may be mechanisms that render extinction context-independent. The AMY is a core hub in fear extinction processing. Data in rodents and humans suggest sustained AMY activity in extinction-impaired individuals, possibly resulting from a failure to engage pro-extinction circuits in cortical areas and subregions of the AMY [reviewed in (Holmes & Singewald, 2013)]. Different AMY subregions and neuronal populations have differential contributions to extinction. The centromedial AMY (CeM) is the major output station of the AMY that drives fear via its connections to the hypothalamus and brainstem regions (Fendt & Fanselow, 1999; LeDoux et al., 1988; Maren, 2001), and its responding is modulated following extinction via intra-amygdala and remote inputs. Extinction training has been shown to cause a rapid reduction of CS-evoked responses of lateral AMY (LA) neurons possibly via depotentiation of thalamic inputs (Duvarci & Pare, 2014). The basal AMY (BA) contains extinction-encoding neurons which drive GABAergic cells in the medial intercalated cell masses (ITC) and neurons in the centrolateral AMY (CeL) to inhibit the CeM and the expression of fear (Duvarci & Pare, 2014; Herry et al., 2008). Following from these initial findings, additional studies involving optogenetic approaches, have shown that a subpopulation of the BA pyramidal neurons which express the Thy1 gene may mark the extinction neuron subpopulation (Jasnow et al., 2013).

It is also becoming apparent that different ITCs are part of a GABAergic feed-forward relay station interconnecting AMY nuclei with distinct networks within the ITCs that are engaged in particular fear stages exerting different influences on extinction (Busti et al., 2011; Duvarci & Pare, 2014; Whittle et al., 2010). Separate neuronal populations in the CeL have been found to have contrasting roles in fear and extinction. CeL-Off neurons (PKC− phenotype) inhibit fear-promoting CeM neurons, while CeL-On (PKC+ phenotype) cells stimulate fear-promoting CeM neurons (Giocchi et al., 2010; Haubensak et al., 2010). Additionally, a very recent finding suggests that a subpopulation of neurons within the CeM – that of the Tac2 peptide-expressing cells – is critically involved in fear learning and fear expression (Andorno et al., 2014).

Another important node within the extinction circuitry is the mPFC, which shares strong interconnections with the HPC and the AMY. Fear extinction recruits the ventral segment of the mPFC, the infralimbic (IL) subdivision [the rodent correlate of the human ventromedial PFC (vmPFC)] which projects to BA extinction and ITC neurons and can thereby inhibit the activity of CeM fear output neurons [reviewed in...]

Fig. 2. Anatomy of fear extinction and expression. Fear extinction and expression rely on neuronal processing in an anatomical circuitry centered on the AMY, mPFC, and HPC. Glutamatergic and GABAergic neurons, among others, are important components of connectivity and regulation of fear. The AMY is critically involved in the expression of aversive (fear) memories. While fear neurons in the BA send excitatory projections directly to the centromedial AMY (CeM) driving expression of fear (right panel), during extinction (left panel), the infralimbic cortex (IL) inhibits CeM output by driving inhibitory ITC neurons. IL inputs might also synapse directly on “extinction” neurons within the BA. “Extinction” neurons can influence activity within the central AMY (CeA) through several routes, possibly by driving inhibitory ITC or CeL (off, PKC−) neurons that limit CeM activity. There is also a BA–CeL pathway contributing to ultimate inhibition of the CeM. The hippocampus is involved in contextual aspects of extinction via its projections to both the IL and the BA, among other brain regions. Hence, inhibitory memories built following extinction are encoded by the AMY and the mPFC and are modulated by the HPC. It is thought that extinction training and exposure therapy produce long-lasting changes in synaptic plasticity and interneuronal communication in this circuitry ultimately reducing fear responses via output stations including the CeM. For further details, the reader is referred to recent reviews of (Duvarci & Pare, 2014; Orsini & Maren, 2012).
3. Neurochemical and molecular substrates of fear extinction

Research is revealing that the activity of a number of specific intracellular signaling cascades [reviewed in (Orsini & Maren, 2012)] carrying biological information from the cell surface to the nucleus, is the key to successful extinction by modulating gene transcription and ultimately promoting synaptic plasticity in extinction-relevant brain regions. The consolidation of extinction memories requires new protein synthesis initiated by molecular signaling cascades within the AMY, HPC and mPFC. The extinction-related molecular signaling and gene expression modulation can differ considerably in these brain areas (Cestari et al., 2014). Key molecules in the amygdala include calcium (Ca2+) influx via NMDA-receptors [in an interaction with α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors] and VGCCs-mediated activation of Ca2+/calmodulin-dependent protein kinase II (CaMKII) and kinases such as Ca2+ /phospholipid-dependent protein kinase (PKC), and Ca2+-dependent protein kinase A (PKA). Once activated, these kinases merge into a common mitogen-activated protein kinase (MAPK)/extracellular regulated kinase (ERK) signaling pathway initiating cAMP response element binding (CREB) phosphorylation and transcription of plasticity proteins [reviewed in (Orsini & Maren, 2012, see Fig. 3 for overview). Aspects of this intracellular signaling seem to be disrupted in deficient extinction, as exemplified by the correlation of impaired extinction with reduced ERK activity in extinction-relevant brain areas (Cannich et al., 2004; Herry et al., 2006; Ishikawa et al., 2012). Similarly, aberrant expression of memory-related genes in these areas is correlated with impaired extinction (Holmes & Singewald, 2013).

To enable selective therapeutic targeting, an important aim in the extinction field is to identify neural mechanisms and intracellular pathways in fear extinction that are different from those involved in fear memory formation. Many of the intracellular mechanisms are similar, however there are also a number of distinct processes and features in fear vs fear extinction learning, including differences in protein synthesis dependence, in brain area- and neuronal population-specific localization and recruitment of signaling pathways, in time course of recruitment, in distinct involvement of certain isoforms of key signaling components (e.g. ERK1 vs ERK2) and resulting gene expression (Cestari et al., 2014; Guedea et al., 2011; Lattal et al., 2006; Tronson et al., 2012). While there are examples of the direct targeting of pharmacologically important components of the aforementioned intracellular signaling, e.g. the MAPK/ERK pathway (Fischer et al., 2007; Herry et al., 2006; Lu et al., 2001) to modulate fear extinction, they were mainly targeted indirectly via membrane receptors e.g. (Cannich et al., 2004; Matsuda et al., 2010) or ion channels (Davis & Bauer, 2012; Ishikawa et al., 2012). These findings have stimulated the hypothesis that boosting specific synaptic plasticity mechanisms by pharmacological means may constitute novel drug targets to promote fear extinction. Indeed, as outlined below, systemic and brain region specific drug studies have identified a number of promising compounds which augment the behavioral and neurobiological effects of fear extinction. The results of these studies are building a framework upon which novel therapeutic interventions to facilitate the efficacy of CBT may be rationally developed and which may ultimately help to enhance symptom remission and recovery.

4. Pharmacologically enhancing extinction

4.1. Serotonergic system

The serotonin (5-HT) system is positioned to modulate the extinction circuitry via ascending 5-HT projections arising from midbrain raphe nuclei that innervate certain brain structures including the AMY (BLA > LA >> CeA), the HPC, the mPFC and the bed nucleus of the stria terminalis [reviewed in (Burgardt & Bauer, 2013)]. The acquisition and expression of conditioned fear increases 5-HT release in the BLA, mPFC and dPAG (Kawahara et al., 1993; Yokoyama et al., 2005; Zanoveli et al., 2009) though possible changes in 5-HT release during acquisition and consolidation of fear extinction have not been reported to the best of our knowledge.

There are 16 5-HT receptor subtypes classified into 7 receptor families (5-HT1–7) (Fig. 4). All 5-HT receptors, excluding the 5-HT4 family—a member of a superfamily of ligand-gated ion channels, are metabotropic, coupled to Gs (5-HT5 and 5-HT7), Gq/11 (5-HT3, 5-HT4 and 5-HT7) or Gq (5-HT2) proteins. To date, research has largely focused on the role of 3 of the 5-HT receptors in fear extinction; namely 5-HT1A, 5-HT2 and 5-HT3 (Table 1, Fig. 4). Selective activation of 5-HT1A (Saito et al., 2013; Wang et al., 2013) or 5-HT2A receptors (Catlow et al., 2013; Zhang et al., 2013) enhances extinction. This might occur via 5-HT1A and 5-HT2A receptors expressed in the mPFC and the LA (Chalmers & Watson, 1991; Cornea-Hebert et al., 1999; Santana et al., 2004). Both of these receptors regulate mPFC and LA excitability via direct activation of pyramidal cells and/or GABAergic interneurons (Illdo-Pelfort et al., 2012; Rainnie, 1999; Stuttmann & LeDoux, 1999). 5HT2A agonists increase presynaptic glutamate release (Aghajanian & Marek, 1999) and increase NMDA receptor sensitivity (Arvanov et al., 1999) and these are mechanisms well characterized as being important in extinction (see Section 4.4 Glutamatergic system).

The roles in extinction of other 5-HT receptors or other components of the 5-HT system, such as melatonin ((Huang et al., 2014) have not been extensively explored. For example, recent work indicating that the 5-HT7 receptor regulates emotional memories (Eriksson et al., 2012), has not been followed up with studies on extinction. However, 5-HT7 receptors have also been associated with fear extinction mechanisms (Table 1). While 5-HT7 antagonists have been found to improve extinction (Park & Williams, 2012), constitutive deletion of 5-HT7 receptors has the opposite effect, possibly due to developmental changes in the 5-HT system (Kondo et al., 2014).

Despite the limited knowledge that we have so far regarding specific 5-HT receptor contributions to extinction, chronic treatment with SSRIs to enhance 5-HT availability is the first-line therapy for many anxiety disorders. Acute SSRI treatment induces anxiogenic effects, which can be prevented with 5-HT2C antagonists in rodents and mimicked by 5-HT2C agonists administered systemically (Bagdy et al., 2001; Salchner & Singewald, 2006) or locally into the BLA (Campbell & Merchant, 2003). Chronic treatment with some SSRIs (e.g., fluoxetine) but not all (e.g., citalopram) enhances extinction in rodents (Table 1) [for a recent review see (Burgardt & Bauer, 2013)], including models of impaired extinction.
Venlafaxine – a combined 5-HT and noradrenaline reuptake inhibitor with weaker affinity to the noradrenaline transporter (Owens et al., 1997) – also improves extinction retrieval and protects against fear reinstatement in rodents (Yang et al., 2012). Chronic fluoxetine treatment does not strengthen fear conditioning or fear expression (Camp et al., 2012) suggesting relative selectivity for extinction. Another clinically relevant attribute of chronic fluoxetine treatment is the protection it provides against the return of fear, as assayed by spontaneous recovery or fear renewal (Deschaux et al., 2011; Deschaux et al., 2013; Karpova et al., 2011). Furthermore, an important molecular mechanism through which fluoxetine produces extinction-associated reductions in fear may be the transformation of adult plasticity mechanisms to a juvenile state (Karpova et al., 2011).

These encouraging preclinical discoveries will hopefully stimulate comprehensive clinical assessment of the efficacy of SSRIs in augmenting CBT in anxiety patients (Table 1A). There have been small-scale clinical trials showing, for example, that paroxetine augments fear reductions (assessed using CAPS scores) and is associated with higher remission rates when combined with CBT in PTSD patients as compared with a placebo-CBT group. An optional treatment maintenance for an additional 12 weeks revealed that symptomatic improvements were long-lasting and consistent although no further improvements could be detected. However, this may have been due to enhanced drop-out
of patients who had remitted in earlier phases of the study (Schneier et al., 2012). More evidence is available in panic disorder (Table 1). There is evidence that chronic SSRI treatment (fluvoxamine, paroxetine) increases CBT-induced fear reductions in panic disorder patients (de Beurs et al., 1995; Oehrberg et al., 1995). However results concerning the extinction-augmenting ability of SSRIs are not consistent as SSRIs, including fluvoxamine, fluoxetine, sertraline and paroxetine, have failed to demonstrate efficacy in primary outcomes [clinical global impression; (Blomhoff et al., 2001; Davidson et al., 2004; Haug et al., 2003; Koszycki et al., 2011; Stein et al., 2000). Despite their common effect of serotonin transporter inhibition, SSRIs are known to differ with regard to their pharmacological properties. Along these lines, some SSRIs (e.g. paroxetine) show additional noradrenaline transporter-inhibiting properties via their metabolites (Owens et al., 1997), while others (e.g. citalopram) show continued selectivity for the serotonergic system (Deupree et al., 2007). In this respect, differential effect sizes of SSRIs are to be expected.

Nevertheless, a number of outstanding questions remain, including that concerning the specificity of SSRIs in augmenting fear extinction as opposed to fear learning (as observed with fluoxetine in preclinical models; see above). In this respect, the finding that prior fear conditioning administration of escitalopram augments extinction learning in healthy volunteers without influencing fear acquisition (Bui et al., 2013) may hint at a potential selectivity of SSRIs to preferentially engage extinction mechanisms. However, this remains to be tested.

In summary, the clinical studies performed so far have demonstrated that SSRIs augmented psychotherapy holds some benefits for panic disorder patients and possibly for PTSD and SAD patients. However, further studies of different SSRIs in larger patient cohorts will be needed, as well as adequate follow-up time to reveal long-term effects, before more definite conclusions can be drawn. Until such studies, as well as studies investigating receptor selective approaches are available, it seems rational to recommend combined treatment involving exposure-based psychotherapy together with antidepressants including SSRIs such as fluoxetine, as there is more evidence supporting than disproving such a strategy at the moment.

### 4.2. Dopaminergic system

An increasing amount of evidence suggests that dopamine (DA) signaling has an important role in extinction mechanisms (Fig. 5) [for a recent detailed review, see (Abraham et al., 2014)]. The DAergic system innervates forebrain extinction circuits through ascending mesocortical/limbic DAergic projections from the ventral tegmental area targeting certain brain regions including the mPFC, HPC and AMY (Pinard et al., 2008; Pinto & Sesack, 2008; Weiner et al., 1991). There
is increased dopamine release in the mPFC during and following extinction training (Hugues et al., 2007) and boosting DAergic signaling with DA precursors or DA-releasing drugs facilitates extinction consolidation (Abraham et al., 2012; Haaker et al., 2013) and can rescue impaired fear extinction in female rats (Rey et al., 2014). Conversely, reducing DAergic transmission by lesioning mesocortical DAergic projection neurons (with 6-OH-DOPA) impairs extinction memory formation (Fernandez Espejo, 2003; Morrow et al., 1999) (see summary in Table 2). More recently, there have been further insights into the mechanisms of how and where DAergic signaling can influence fear extinction have been made. These studies (see Table 2 and below) have revealed a complex two-pronged feature of DAergic signaling that can (i) gate the expression of fear, and (ii) influence fear extinction consolidation mechanisms.

Dopamine receptors are metabotropic and in the classical view, coupled to either Gs (increased cAMP, D1-like) or Gi/0 (decreased cAMP, D2-like) proteins. However, DAergic signals can also be transduced via Phospholplase C (PLC) activation (enhanced DAG, IP3) mediated by D1-like Go proteins, by D2-like Gi/0 subunits or by receptors forming D1/D2 heteromers (Felder et al., 1989; Lee et al., 2004) [for review see (Abraham et al., 2014)]. The D1-class receptor signaling modulates the excitability of BLA parvalbumin-positive interneurons (Bissiere et al., 2003; Kroner et al., 2005; Lorentz et al., 2004) and ITCs (Manko et al., 2011). In concert with D2-like receptors in the CeL/CeC (Perez de la Mora et al., 2012), DAergic signaling exerts tight control over CeM-mediated expression of fear. To date, however, pharmacological manipulation of DAergic signaling in the AMY has produced conflicting results with regard to extinction (Fiorenza et al., 2012; Hikind & Maroun, 2008) (see Table 2).

Preclinical work has aimed to clarify the role of DA receptor activity in extinction. However, research has been largely complicated by the lack of receptor subtype-selective compounds, leaving many investigations reliant on drugs acting on D1-like (D1, D5) and D2-like (D2, D3, D4) receptor subfamilies. One reliable finding, however, is that DA antagonism (see Table 2) significantly in the mPFC impairs the retrieval of extinction memories (Hikind & Maroun, 2008; Mueller et al., 2010; Pfeiffer & Fendt, 2006). In this case, DAergic effects on glutamatergic NMDA receptor signaling may be involved. As discussed below (Section 4.4 Glutamatergic system), NMDA receptor-mediated burst activity of mPFC neurons is associated with stabilization of extinction memories (Burgos-Robles et al., 2007) and D1-like receptor agonists facilitate NMDA receptor currents via Gs mediated increases in adenylyl cyclase (AC) activity (Snyder et al., 1998). There is also the possibility that D2 receptor mechanisms in the PFC support extinction, as (Mueller et al., 2010) showed that a pre-extinction IL injection of the D2 antagonist raclopride impaired the consolidation of extinction memory. However, following systemic administration of the (less selective) D2 receptor antagonist sulpiride, accelerated extinction was found (Ponnusamy et al., 2005).

In summary, the results of preclinical studies do not yet fully clarify the specific DA receptor or receptor class mediating the extinction augmenting effect of enhanced dopaminergic signaling. Work still needs to be done on the use of receptor-selective agonists and antagonists, as...
well as biased ligands (showing functional selectivity for either AC or PLC signal transduction pathways) to reveal receptor-specific functions and their modulation of downstream signaling cascades in extinction. Pharmacologically increasing the DA tone promotes extinction. L-DOPA administration following extinction training enhances fear extinction in mice and healthy humans, rendering extinction resistant to fear renewal, reinstatement and spontaneous recovery (Haaker et al., 2013). L-DOPA is a precursor for all catecholamines, but preferentially enhances dopaminergic turnover in the frontal cortex (Dayan & Finberg, 2003), eliciting D1- and D2-like receptor responses (Trugman et al., 1991). Indeed, L-DOPA augments extinction-related neural activity in the mPFC (IL) of mice and increases mPFC functional coupling (Mueller et al., 2008) and successful fear extinction is associated with enhanced extracellular levels of noradrenaline in the mPFC (Hugues et al., 2007). Boosting noradrenaline levels, via administration of either noradrenaline itself (Merlo & Izquierdo, 1967) or of compounds such as yohimbine (see below) or methylphenidate (Abraham et al., 2012) enhances fear extinction (Table 3). There is evidence from preclinical studies that activating α2-adrenoceptors, one target receptor of noradrenaline, can also facilitate fear extinction (Table 3) [for recent review, see (Fitzgerald et al., 2014a)]. Conversely, depleting central noradrenaline or lesioning ascending noradrenaline projections from the locus coeruleus, impairs extinction, as does systemic alpha1 or β1 adrenoceptor blockade (Table 3) [reviewed in (Mueller & Cahill, 2010)].

### 4.3. Noradrenergic system

The noradrenergic system is crucial for both the formation and the maintenance of fear memories, as well as for extinction memories (reviewed in (Holmes & Quirk, 2010; Mueller & Cahill, 2010)). Noradrenaline can enhance neuronal excitability in extinction-relevant brain regions such as the IL (Mueller et al., 2008) and successful fear extinction is associated with enhanced extracellular levels of noradrenaline in the mPFC (Hugues et al., 2007). Boosting noradrenaline levels, via administration of either noradrenaline itself (Merlo & Izquierdo, 1967) or of compounds such as yohimbine (see below) or methylphenidate (Abraham et al., 2012) enhances fear extinction (Table 3). There is evidence from preclinical studies that activating β-adrenoceptors, one target receptor of noradrenaline, can also facilitate fear extinction (Table 3) [for recent review, see (Fitzgerald et al., 2014a)]. Conversely, depleting central noradrenaline or lesioning ascending noradrenaline projections from the locus coeruleus, impairs extinction, as does systemic alpha1 or β1 adrenoceptor blockade (Table 3) [reviewed in (Mueller & Cahill, 2010)].

There has been interest in the clinical utility of targeting noradrenergic mechanisms to augment extinction. Here, we focus on α2-adrenoceptors given the recent clinical findings supporting the utility of α2-adrenoceptor antagonists in improving extinction. Based on results of rodent studies (Cain et al., 2004), two small clinical studies (Powers et al., 2003; Bui et al., 2013) revealed significant and sustained improvements in skin conductance responses, with patients showing no significant differences in primary CGI outcome. Secondary outcome: Higher proportion of panic-free patients in paroxetine-CBT group.

### Table 1A

| Disorder | Study design | Outcome (compared to placebo control group) | Reference |
|----------|-------------|---------------------------------------------|-----------|
| Healthy volunteers | 14 days post escitalopram pretreatment; fear conditioning paradigm | Accelerated extinction learning (skin conductance responses) after extinction, as does systemic alpha1 or β1 adrenoceptor blockade (Table 3) [reviewed in (Mueller & Cahill, 2010)]. | (Bui et al., 2013) |
| Panic disorder with/without agoraphobia | Fluvoxamine or placebo followed by exposure therapy; psychological panic management followed by exposure therapy or exposure therapy alone | Self-reported measures, All treatments effective, however fluvoxamine plus CBT superior to all other treatments | (de Beurs et al., 1995) |
| Panic disorder with/without agoraphobia | 12 weeks paroxetine or placebo plus CBT | Reduced number of panic attacks in paroxetine/CBT group | (Oehrberg et al., 1995) |
| Panic disorder with/without agoraphobia | 10 weeks of paroxetine or placebo plus CBT in week 5 and 7 | No significant difference in primary CGI outcome | (Stein et al., 2000) |
| Panic disorder with/without agoraphobia | 12 weeks fluvoxamine or placebo with or without CBT | All groups improved (also placebo without CBT) | (Sharp et al., 1997) |
| Panic disorder with/without agoraphobia | 12 weeks of sertraline or placebo treatment plus self-administered CBT or no CBT | Reduced anticipatory anxiety in sertraline plus self-administered CBT | (Koszycki et al., 2011) |
| Social anxiety disorder | 24 weeks of sertraline/placebo with or without exposure therapy (8 sessions in the first 12 weeks of treatment) | No significant improvements in CGI. | (Blomhoff et al., 2001) |
| Social anxiety disorder | Follow-up study of (Blomhoff et al., 2001) Assessment of long-term effects 28 weeks after cessation of medical treatment | Exposure therapy alone (without placebo or sertraline) showed a further improvement in CAPS 28 weeks after treatment cessation, however only reached improvement levels comparable with those of the sertraline alone group after the initial 24 weeks | (Haug et al., 2003) |
| Social anxiety disorder | 14 weeks of fluoxetine or placebo plus weekly CBT or no CBT CBT consisted of group treatment combining in vivo exposure, cognitive restructuring and social skills training | All treatments were superior to placebo (without CBT) but no differences between groups themselves | (Davidson et al., 2004) |
| PTSD | 10 exposure therapy sessions (1×/week) plus paroxetine CR Optional 12 weeks of maintenance treatment | Greater CAPS improvement in paroxetine group vs placebo after 10 weeks | (Schneier et al., 2012) |

CR...controlled release.
et al., 2009; Smits et al., 2014) have shown that combining yohimbine with CBT can reduce anxiety in patients with social anxiety disorder and claustrophobic fear (Table 3A). A third study found no CBT-augmenting benefit of yohimbine in patients with fear of flying; however, CBT may have exerted a ‘ceiling effect’ that occluded the drug’s effect (Meyerbroeker et al., 2012).

In rodents, blocking α2-adrenoceptors (e.g., with yohimbine) that function as autoreceptors on locus coeruleus neurons increases locus coeruleus activity and noradrenaline release in terminal regions (Singewald & Philippu, 1998). Yohimbine is anxiogenic and increases neuronal activity in widespread brain areas, including extinction-relevant areas such as the amygdala, mPFC and HPC (Singewald et al., 2003). Downstream targets of yohimbine-induced noradrenaline release include β1-adrenoceptors, and increasing activity at these receptors in locus coeruleus terminal regions (mPFC) enhances extinction (Do-Monte et al., 2010b), possibly via facilitating extinction-relevant long-term potentiation (Gelinas & Nguyen, 2005) in a BDNF-dependent manner (Furini et al., 2010). Yohimbine facilitates extinction in rodents, including extinction-impaired subjects (Hefner et al., 2008, 2008), though a more selective α2-adrenoceptor, atipamezole, does not (Table 3) [for further discussion, see (Holmes & Quirk, 2010)]. Yohimbine also protects against spontaneous fear recovery, in a manner similar to the effect of a noradrenaline uptake inhibitor, atomoxetine (Janak & Corbit, 2011). However, yohimbine reduces fear only in the extinction context in which the drug was administered (Morris & Bouton, 2007), which would be a limitation of the clinical goal of achieving context-independent reductions in fear. Thus, other strategies may be needed, including combination strategies where drugs which do produce context-independent extinction, for example targeting HDACs (see Section 4.10 Epigenetics), are combined with yohimbine.

4.4. Glutamatergic system

Glutamatergic signaling plays a crucial role in synaptic plasticity and many forms of learning and memory, including fear extinction (see Fig. 6 for an overview). Fast excitatory glutamatergic signaling is mediated by ionotropic receptors (NMDA and AMPA), and slower signaling by metabotropic (mGluR1–8) receptors. Group I (mGluR1 and
mGlur5 receptors are mainly expressed postsynaptically and coupled to Gq proteins enhancing phospholipase C (PLC) activity. Group II (mGlur2, mGlur3) and Group III (mGlur4, mGlur6–mGlur8) are both coupled to Gi proteins inhibiting adenylate cyclase (AC) activity and expressed on pre- as well as postsynaptic sites (Willard & Kochenbour, 2013).

Pharmacological potentiation of AMPA receptor activation (by PEPA) facilitates extinction learning and retrieval (Yamada et al., 2009, 2011; Zushida et al., 2007), although it is ineffective in severely extinction-impaired subjects (Whittle et al., 2013). Studies using localized infusion of AMPA potentiators (Zushida et al., 2007) and blockers (Falls et al., 1992; Milton et al., 2013; Zimmerman & Maren, 2010) coupled with electrophysiological recordings suggest that AMPA-mediated effects on extinction are localized to the mPCF (Zushida et al., 2007) see Table 4, for recent detailed reviews (Bukalo et al., 2014; Myers et al., 2011).

The part played in extinction by the metabotropic glutamate receptors mGlur1, mGlur5 and mGlur7 has also been evaluated. Antagonizing mGlur1 (which is expressed on ITC innervating neurons (Busti et al., 2011) with CPPCOEt (Kim et al., 2007) and antagonizing mGlur5 receptors in the IL (Sepulveda-Oreno et al., 2013) have been shown to impair extension. Furthermore, gene mutations resulting in deficits in mGlur5 (Xu et al., 2009) or mGlur7 (Callaerts-Vegh et al., 2006; Fendt et al., 2008; Goddyn et al., 2008) impair extinction learning, while selective activation (Dobi et al., 2013; Fendt et al., 2008; Morawska & Fendt, 2012; Rodrigues et al., 2002; Siegel et al., 2008; Toth et al., 2012b; Whittle et al., 2013) improves extinction [see Table 4 for summary and (Bukalo et al., 2014; Myers et al., 2011) for more details].

The most extensively studied glutamate receptor in relation to fear extinction is the NMDA receptor (NMDAR). As shown in Table 4, systemic or local administration of NMDAR antagonists into the BLA or mPCF produces extinction deficits [reviewed in (Myers et al., 2011)]. Due to the potential for major side-effects (e.g. excitotoxicity) a general enhancement of NMDAR transmission is clinically undesirable and more subtle modulation of NMDAR signaling is required. NMDARs are specialized voltage-dependent, ligand-gated ion channels that are expressed as heterotetramers formed by GluN1 in combination with GluN2 or GluN3 subunits. To date, eight different splice variants of the GluN1 subunit, four distinctive GluN2 (GluN2A–D) subunits and two GluN3 (GluN3A, B) subunits have been identified [for review see (Paoletti et al., 2013)]. Activation of NMDARs requires L-glutamate binding to GluN1 subunits, L-glutamate or D-serine binding to GluN2 subunits and membrane depolarization relocating the channel pore-blocking Mg2+ and Zn2+ ions.

The repertoire of NMDAR subunits permits the assembly of a range of NMDARs with varying dissociable signaling properties. Systemic (Dalton et al., 2008; Dalton et al., 2012; Leaderbrand et al., 2014; Sotres-Bayon et al., 2007) or localized BLA (Laurent & Westbrook, 2008; Sotres-Bayon et al., 2007; Sotres-Bayon et al., 2009) or mPCF (Laurent & Westbrook, 2008; Sotres-Bayon et al., 2009) inhibition of GluN2p-containing NMDARs (via ifenprodil or Ro25-6981) disrupts extinction. Conversely, GluN2B overexpression...
enhances extinction (Tang et al., 1999). Recently, facilitated signaling of GluN2C/D-containing NMDARs in the BLA (via CIQ infusion) was also demonstrated to enhance extinction (Ogden et al., 2014). The contribution of other subunits, such as GluN2A, remains to be determined.

The organization of NMDARs provides additional druggable pharmacological targets. GluN1 and GluN2 subunits are endowed with binding sites for positive and negative allosteric regulations that enable fine-tuning of NMDAR activity. The GluN1 subunit contains an inhibitory H+ binding site rendering GluN2B/D-containing NMDARs in particular sensitive to local pH changes, as well as to endogenous molecules with redox potential (Banke et al., 2005). The GluNR2 subunit is responsive to allosteric modulation of NMDAR signaling by polyamines such as spermidine. Spermidine-mediated activation of GluN2B signaling facilitates the consolidation of extinction (Gomes et al., 2010; Guerra et al., 2006), while aracine, an antagonistic polyamine, disrupts extinction (Gomes et al., 2010). A GluN2 allosteric binding site – still uninvestigated in the field of fear extinction – enables modulation of NMDAR signaling by neurosteroids (I.a. allopregnanolone) (Irwin et al., 1994).

Nuanced pharmacological modulation of NMDARs can also be achieved by targeting the Zn2+ binding domain found on GluN2 subunits. Zn2+-binding to this modular site mainly inhibits GluN2A-containing NMDARs, by reducing NMDAR channel open probability (Paoletti et al., 1997). Brain Zn2+ signaling which, in addition to the NMDAR binding site, acts via other sites including the GPR39 Zn-sensing receptor (Holst et al., 2007), has been shown to exert both enhancing and attenuating effects on learning and memory via complex interactions with neurotransmitters and synaptic plasticity mechanisms (reviewed in Takeda & Tamano, 2014). Dietary Zn2+ supplementation impairs extinction (Railey et al., 2010), while dietary-induced reduction of brain Zn2+ levels rescues deficient extinction (Whittle et al., 2010). Although these effects may be mediated by Zn2+ modulation of NMDAR signaling, other actions cannot be excluded (e.g., see discussion on HDAC inhibition in Section 4.10: Epigenetics). The same is true for another NMDAR-modulating ion, Mg2+, which has also been shown to facilitate extinction when globally enhanced in the brain (Abumaria et al., 2011; Mickley et al., 2013).

The most extensively investigated allosteric modulation of NMDARs is mediated via the γ-glycine binding site on the GluN1 subunit. As shown in Table 4, systemic administration of ligands on this binding site, n-serine or n-cycloserine (DCS), augments extinction consolidation [see (Bukalo et al., 2014; Myers et al., 2011) for recent detailed reviews]. There is evidence (although not consistent) that DCS may promote generalization of the extinction effect (Lederwood et al., 2005; Vervliet, 2008) that is, to other cues (e.g., odors, sounds, visual stimuli) that acquired fear during a more complex conditioning situation. Generalized extinction could be of potential clinical benefit, provided that cues that trigger adaptive reactions are not affected. However, DCS administration outside the consolidation window can limit the drug's effectiveness. When extinction or exposure sessions are too short or yield insufficient fear induction, DCS can strengthen (re-)-consolidation of the original fear memory. Along these lines, the extinction augmenting effects of DCS require the subject to show at least some ability to extinguish (Bolkan & Lattal, 2014; Bouton et al., 2008; Hefner et al., 2008; Smits et al., 2013b; Tomilenko & Dubrovina, 2007; Weber et al., 2007; Whittle et al., 2013). Finally, chronic DCS treatment leads to loss of effects on extinction, possibly via NMDAR desensitization (Parnas et al., 2005; Quartermain et al., 1994). It is of note, that chronic treatment with antidepressants also disrupts the facilitating effects of DCS on extinction, possibly by interfering with NMDAR function (Werner-Seidler & Richardson, 2007).

Notwithstanding these caveats, DCS represents one of the best examples of translational research paving the way for novel anxiety treatments. DCS augmentation of CBT has been clinically studied in various anxiety disorders (Table 4A) including among pediatric patients [see (Hofmann et al., 2013a) for a recent review]. To summarize the current clinical data, DCS-augmented CBT shows advantages in PTSD (de Kleine et al., 2012; Difede et al., 2014; Scheerling & Weems, 2014), specific phobia (Guastella et al., 2007; Nave et al., 2012; Ressler et al., 2004), SAD (Guastella et al., 2008; Hofmann et al., 2006; Hofmann et al., 2013b; Smits et al., 2013c) and OCD (Chasson et al., 2010; Farrell et al., 2013; Kushner et al., 2007; Storch et al., 2010; Wilhelm et al., 2008) but see (Mataix-Cols et al., 2014). Panic disorder patients (Otto et al., 2010c; Siegmund et al., 2011) also show faster symptom alleviation after DCS-augmented CBT. Reflecting the preclinical findings, negative outcomes were associated with short CBT sessions (Utz et al., 2012), insufficient within-session fear inhibition (Smits et al., 2013b; Tart et al., 2013), chronic DCS treatment (Heresco-Levy et al., 2002), DCS administration outside the therapeutic temporal window (Storch et al., 2007) and subclinical levels of fear [(Guastella et al., 2007; Gutner et al., 2012) but see (Kuriyama et al., 2011)]. If these factors are properly attended to [see above, reviewed in (Hofmann, 2014)], the evidence supports the adjunctive treatment of CBT with DCS for a range of anxiety disorders.

4.5. γ-Aminobutyric acid (GABA)

GABA is the main inhibitory neurotransmitter in the adult mammalian brain. Because extinction largely reflects the promotion of active inhibitory processes, it is not surprising that GABA serves as an important source of such inhibition. GABA signaling occurs largely via binding to ionotropic GABAA and metabotropic GABAB receptors (former GABAC receptors have been reclassified and are now termed GABAa rho subclass receptors). GABAA receptors are pentameric, ligand-gated CI− channels which, upon activation induce hyperpolarization of cells and hence reduce neuronal excitability. Seven classes of GABAA subunits (α1-6, β1-3, γ1-3, π1-3, δ, ε, ) have been identified so far, permitting the assembly of highly variable GABAA receptors with distinct functions and signaling properties. Furthermore, several allosteric binding sites, in addition to the GABA binding pocket located at the α3-β-subunit interface, allow nuanced modifications of GABAa receptor signaling. The allosteric benzodiazepine (BDZ) binding site is localized at the interface of α- and γ-subunits and when activated, facilitates GABAergic transmission by promoting GABA binding to the receptor and increasing the opening probability of the CI− channel. The binding sites of other allosteric modulators of GABAa signaling including barbiturates and neurosteroids among others are not yet fully characterized [see Fig. 7 and (Gunn et al., 2014; Makkar et al., 2010)].
In preclinical studies, the robust inhibitory effect of full GABA<sub>A</sub> receptor agonists at the GABA<sub>A</sub> binding site (α/β-subunit interface) (e.g. muscimol), is often used to inactivate a brain region to clarify its participation in certain functions including extinction. Muscimol-injections into the BLA [Laurent & Westbrook, 2008; Laurent et al., 2008; Sierra-Mercado et al., 2011], mPFC/IL [Laurent & Westbrook, 2008, 2009; Sierra-Mercado et al., 2011] and HPC [Corcoran et al., 2005; Sierra-Mercado et al., 2011], see Table 5 for a summary, disrupt extinction learning and memory, supporting the crucial involvement of these areas. However, muscimol infusion into the BLA and IL facilitates extinction in some studies [Akiwar et al., 2006].

There is solid evidence that fear extinction learning is associated with upregulation of GABAergic markers in the AMY, underscoring the importance of a certain level of GABAergic signaling in this brain area for the successful formation of extinction memories. For example, levels of gephyrin, a clustering protein facilitating postsynaptic scaffolding of GABA<sub>A</sub> receptors, were enhanced 2 h following extinction training along with enhanced surface expression of GABA<sub>A</sub> receptors in these areas. However, muscimol infusion into the BLA and IL facilitates extinction in some studies [Akiwar et al., 2006].

Table 3
Noradrenergic (NE) signaling in fear extinction (preclinical studies).

| Drug/manipulation          | Extinction learning | Extinction retrieval | Longterm extinction | Route | Reference                  |
|----------------------------|---------------------|----------------------|----------------------|-------|---------------------------|
| Enhancing noradrenergic signaling |                     |                      |                      |       |                           |
| Noradrenaline              | ns                  | +                    | ns                   | icv   | (Merlo and Izquierdo, 1967) |
| Methylphenidate            | +                   | +                    | ns                   | ip    | (Abraham et al., 2012)    |
| Noradrenaline              | ns                  | +                    | ns                   | BLA   | (Berlau and McGaugh, 2006) |
| Atomoxetine (NE reuptake inhibitor) | ns                  | No effect            | ns                   | BLA   | (Fiorenza et al., 2012)   |
| Isoproterenol (β-agon)     | No effect           | (+)                  | ns                   | ip/subchr (chonic) | (Do-Monte et al., 2010b) |
| Yohimbine (α2)             | +                   | No effect            | ns                   | icv   | (Do-Monte et al., 2010b)  |
| Atipamezole (α2)           | ns                  | No effect            | (+)                  | sc    | (Davis et al., 2008)      |

Inhibiting noradrenergic signaling

| Prazosin (α1)             | ns                  | No effect            | ns                   | ip/subchr (chonic) | (Do-Monte et al., 2010a) |
| Propanolol (β-ant)        | ns                  | No effect            | ns                   | sc    | (Do-Monte et al., 2010a)  |
| Sotalol (β-ant)           | ns                  | No effect            | ns                   | mPFC   | (Do-Monte et al., 2010a)  |
| Timolol (β-ant)           | ns                  | No effect            | ns                   | sc    | (Rodriguez-Romaguera et al., 2009) |
| Atenolol (β-ant)          | ns                  | No effect            | ns                   | mPFC   | (Do-Monte et al., 2010a)  |
| Methylphenidate           | (+)                 | (+)                  | ns                   | sc    | (Do-Monte et al., 2010a)  |
| Yohimbine (α2)            | ns                  | No effect            | (+)                  | sc    | (Do-Monte et al., 2010a)  |
| Atipamezole (α2)          | ns                  | No effect            | (+)                  | mPFC   | (Do-Monte et al., 2010a)  |

1 Drug administration following extinction training; 2 spaced CS extinction, US = 0.7mA; 3 spaced CS extinction when US = 0.4mA; no effect when US = 0.7mA; 4 cocaine-conditioned place preference; 5 systemic (Janak and Corbit, 2011); 6 +, Improved; –, impaired; (+) or (–), only minor effects; ip, intraperitoneal injection; injection; sc, subcutaneous injection; icv, intracerebroventricular injection; ns, not studied; HPC, hippocampal administration; BLA, intra-basolateral amygdala administration; IL, infralimbic cortex; PL, prelimbic cortex; CA1, cornu ammonis 1; Ren, Fear renewal; SR, spontaneous recovery; Re-in, reinstatement; ag, agonist; ant, antagonist, KO, knock-out; #, reduced fear expression at start of extinction training.

Table 3A
Yohimbine combined with CBT in small-scale clinical trials.

| Disorder            | Study design                  | Outcome (compared to placebo control group) | Reference                  |
|---------------------|-------------------------------|--------------------------------------------|----------------------------|
| Social anxiety disorder (SAD) | 4 sessions consisting of yohimbine being administered prior to a CBT + session | Yohimbine augmented CBT-induced improvement in SAD (self-report measures; 21 day follow-up) | (Smits et al., 2014) |
| Claustrophobic fear | 1 single oral yohimbine dose prior to an exposure + session | Yohimbine augment CBT-induced reduced fear of enclosed spaces (7 day follow-up) | (Powers et al., 2009) |
| Fear of flying      | 2 single oral yohimbine prior to VRET session | No fear augmenting effect of yohimbine (Meyerbroeker et al., 2012) | (Meyerbroeker et al., 2012) |
the BLA (Chhatwal et al., 2005a). Extinction learning also increases GABA<sub>A</sub> receptor α<sub>2</sub>- and β<sub>2</sub> subunit mRNA and levels of the GABA-synthesizing enzyme glutamate decarboxylase (GAD) and the GABA transporter GAT1 (Heldt & Ressler, 2007). Further demonstrating the importance of GABAergic signaling to extinction, fear extinction learning is disrupted by mutation-induced deficits in the activity-dependent GAD isoform GAD65 (Sangha et al., 2009), and by viral knock-down of the constitutive GAD isoform GAD67 (Heldt et al., 2012). While these findings suggest that upregulation of GABAergic signaling supports extinction, GABA<sub>A</sub> receptor antagonists such as picROTOXIN or bicuculline can enhance extinction memories when administered systemically (McGaugh et al., 1990), or when infused into the BLA (Berlau & McGaugh, 2006) or specific (fear promoting) areas (PL) of the mPFC (Fitzgerald et al., 2014b; Thompson et al., 2010). The reason for these apparent contradictions is not yet clear.

Preliminary findings from studies using genetically modified mice suggest that the development of subunit-selective GABA<sub>A</sub> receptor drugs could be a promising way to more distinctly utilize GABAergic mechanisms facilitating extinction. For example, there is reduced mRNA expression of α<sub>5</sub> and γ<sub>1</sub> subunits in the AMY of a mouse model of enhanced anxiety (Tasan et al., 2011) displaying impaired extinction (Yen et al., 2012). In addition, deletion of α<sub>5</sub>-containing GABA<sub>A</sub> receptors in the HPC is sufficient to impair fear extinction recall (Yee et al., 2004) while global deletion of the GABA<sub>A</sub> α<sub>3</sub> subunit produces modest improvements in extinction learning (Fiorelli et al., 2008).

An additional possible explanation for some paradoxical findings concerning GABA regulation and AMY function in extinction is the fact that most of the prior studies have performed pharmacological or genetic manipulations that affect either the entire brain, just the AMY, or even only AMY subregions. Despite this, we now know that GABA...
### Table 4
Glutamatergic signaling in fear extinction (preclinical studies).

| Drug/manipulation          | Extinction training | Extinction retrieval | Longterm extinction | Route | Reference                                      |
|----------------------------|---------------------|----------------------|----------------------|-------|------------------------------------------------|
| **Facilitating AMPA signaling** |                     |                       |                      |       |                                                |
| PEPA (AMPA potentiator)    | No effect           | No effect            | ns                   | ip    | (Whittle et al., 2013)                         |
|                            |                     | +                    | ns                   | ip    | (Yamada et al., 2011)                         |
|                            | +                   | +                   | ns                   | ip    | (Yamada et al., 2009)                         |
|                            | +                   | +                   | + (Re-in)            | ip    | (Zushida et al., 2007)                        |
| PEPA + NBQX (AMPA ant)     | No effect           | No effect            | ns                   | ip    | (Zushida et al., 2007)                        |
|                            | (+)                 | (+)                 | ns                   | PL    | (Zushida et al., 2007)                        |
|                            | +                   | +                   | ns                   | BLA/ceA | (Zushida et al., 2007)                        |
| **Inhibiting AMPA signaling** |                     |                       |                      |       |                                                |
| CNQX (AMPA ant)            | ns                  | No effect            | ns                   | BLA   | (Falls et al. 1992; Lin et al. 2003b; Zimmerman and Maren 2010) |
| **Facilitating NMDA signaling** |                     |                       |                      |       |                                                |
| D-Cycloserine              | ns                  | +                   | ns                   | ip    | (Walker et al., 2002)                         |
|                            | +                   | +                   | ns                   | Systemic | (Ledgerwood et al., 2003)                     |
|                            | +                   | +                   | + (Re-in)            | Systemic | (Ledgerwood et al., 2004)                     |
|                            | +                   | +                   | ns                   | Systemic | (Ledgerwood et al., 2005)                     |
|                            | +                   | +                   | ns                   | ip    | (Parnas et al., 2005)                         |
|                            | +                   | +                   | ns                   | ip    | (Yang and Lu, 2005)                           |
|                            | +                   | +                   | ns                   | ip    | (Lee et al., 2006)                            |
|                            | +                   | +                   | ns                   | No effect | sc | (Woods and Bouton 2006)                       |
|                            | +                   | +                   | ns                   | Systemic | (Werner-Seidler and Richardson, 2007)         |
|                            | +                   | +                   | + (Re-in)            | Systemic | (Weber et al., 2007)                         |
|                            | +                   | +                   | ns                   | ip    | (Yang et al., 2007)                           |
|                            | +                   | +                   | ns                   | ip    | (Yang et al., 2007)                           |
|                            | +                   | +                   | ns                   | ip    | (Hefner et al., 2008)                         |
|                            | +                   | +                   | ns                   | po    | (Yamamoto et al. 2008)                       |
|                            | +                   | +                   | ns                   | ip    | (Matsumoto et al., 2008)                      |
|                            | +                   | +                   | ns                   | ip    | (Silvestri and Root, 2008)                    |
|                            | +                   | +                   | ns                   | Systemic | (Langton and Richardson, 2008)                |
|                            | +                   | +                   | No effect            | sc    | (Bouton et al., 2008)                         |
|                            | +                   | +                   | ns                   | Systemic | (Lin et al., 2010)                            |
|                            | +                   | +                   | + (Re-in)            | ns    | (Whittle et al., 2009)                        |
|                            | +                   | +                   | No effect            | Systemic | (Langton and Richardson, 2010)                |
|                            | +                   | +                   | ns                   | ip    | (Yamada et al., 2011)                         |
|                            | +                   | +                   | ns                   | ip    | (Toth et al., 2012b)                          |
|                            | +                   | +                   | ns                   | ip    | (Bai et al., 2014)                            |
|                            | +                   | +                   | + (Re-in)            | ns    | (Whittle et al., 2013)                        |
|                            | +                   | +                   | ns                   | ip    | (Walker et al., 2002)                         |
|                            | +                   | +                   | ns                   | ip    | (Ledgerwood et al., 2003)                     |
|                            | +                   | +                   | ns                   | ip    | (Lee et al., 2006)                            |
|                            | +                   | +                   | ns                   | ip    | (Lee et al., 2008)                            |
|                            | +                   | +                   | ns                   | BLA   | (Walker et al., 2002)                         |
|                            | +                   | +                   | ns                   | BLA   | (Lee et al., 2006)                            |
|                            | +                   | +                   | ns                   | BLA   | (Lee et al., 2008)                            |
|                            | +                   | +                   | ns                   | BLA   | (Lee et al., 2006)                            |
|                            | +                   | +                   | No effect            | BLA   | (Boklan & Lattal, 2014)                       |
|                            | +                   | +                   | ns                   | HPC   | (Boklan & Lattal, 2014)                       |
|                            | +                   | +                   | +                   | HPC   | (Ben et al., 2010)                            |
|                            | +                   | +                   | ns                   | HPC   | (Gomes et al., 2010)                          |
| **Spermidine**             |                     |                       |                      |       |                                                |
| **Inhibiting NMDA signaling** |                     |                       |                      |       |                                                |
| MK-801 (non-competitive NMDA ant) | ns                  | ns                   | Systemic             | Systemic | (Baker & Azorlosa, 1996); (Storsve et al., 2010) |
|                            | ns                  | ns                   | + (Re-in)            | sc    | (Johnson et al., 2000)                        |
|                            | ns                  | −                   | ns                   | ip    | (Lee et al., 2006); (Liu et al., 2009)         |
|                            | ns                  | −                   | Systemic             | Systemic | (Langton et al., 2007); (Chan & McNally, 2009) |
| **CPP (competitive NMDA ant)** | No effect           | −                   | ns                   | ip    | (Santini et al., 2001); (Sotres-Bayon et al., 2007) |
|                            | No effect           | −                   | ns                   | MPC   | (Burgos-Robles et al., 2007)                   |
|                            | −                   | −                   | ns                   | MPC   | (Burgos-Robles et al., 2007)                   |
|                            | −                   | −                   | ns                   | BLA   | (Parsons et al., 2010)                        |
|                            | −                   | −                   | ns                   | BLA   | (Falls et al., 1992)                          |
|                            | −                   | −                   | ns                   | BLA   | (Lee & Kim, 1998)                             |
|                            | −                   | −                   | ns                   | BLA   | (Lin et al., 2001a); (Laurent et al., 2008); (Fiorenza et al., 2012) |
|                            | No effect           | −                   | ns                   | BLA   | (Lin et al., 2003a); (Zimmerman & Maren, 2010) |
|                            | −                   | −                   | ns                   | CA1   | (Szapiro et al., 2003)                        |
|                            | −                   | −                   | ns                   | CA1   | (Fiorenza et al., 2012)                       |
|                            | −                   | −                   | ns                   | MPC   | (Fiorenza et al., 2012)                       |
receptors are found on most of the neuronal cell types and thus even a subregion-specific modulation is likely to affect many different cell populations which may have opposing functional effects. One recent attempt to address this complexity involved the use of an inducible GABA B1b knockout strategy limited only to the corticotropin-releasing hormone containing neuronal population (Gafford et al., 2012), which revealed relatively specific effects, including enhanced anxiety and extinction deficits. It is likely that future studies aimed at cell-type specific manipulations will help address the vast complexity of the many different cell types combined with the large number of different GABA receptor populations.

As mentioned above, the multiple different binding sites found within GABA A receptors can be used to elicit a more nuanced modulation of GABA A receptor activity. Positive allosteric modulation of GABA A receptor activity via BZD, represents the most widely described drug-based strategy for acute treatment of anxiety states (Macaluso et al., 2010). The prevalence of BZD use among patients suffering from mental disorders is high, and is estimated to range between 40 and 70% of these patients (Clark et al., 2004) However anxiolytic treatment with BZDs is recommended only for short periods of time, due to several disadvantages including its sedative effects, the development of dependence and difficulties with discontinuing chronic treatment (Otto et al., 2002). The sedating and calming effects of BZDs might in fact interfere with extinction possibly via reduced arousal and decreased release of neurotransmitters (e.g. noradrenaline) and stress hormones (e.g. glucocorticoids) that support extinction ([Bentz et al., 2010], see Section 4.8 Glucocorticoids for details). BZDs may also render extinction memories dependent on the drug-induced internal anxiolytic-state (state dependent learning), which upon BZD discontinuation may no longer be accessible. These concerns are borne out by preclinical data showing that systemic (Bouton et al., 1990; Bustos et al., 2009; Goldman, 1977; Hart et al., 2009, 2010, 2014; Pereira et al., 1989) or intra-BLA (Hart et al., 2009, 2010) BZD administration impairs extinction (Table 5), presumably also through state-dependent mechanisms. Systemic treatment with a BZD inverse agonist (Harris & Westbrook, 1998; Kim & Richardson, 2007, 2009) has similar effects (see Table 5 for a summary). Hence, combining BZD with CBT needs careful consideration. As well as further development of BZDs targeting specific subunits of the GABA A receptor [reviewed in (Griebel & Holmes, 2013)], which provides some hope, there is preliminary evidence that putative novel anxiolytics that do not impair extinction learning [e.g. neuropeptide S; (Slattery et al., in press)] can be de-
receptors (GABA_{B1a,b} knock-out) impairs extinction learning (Jacobson et al., 2006), pharmacological GABA_{B} receptor antagonism does not (Heaney et al., 2012; Sweeney et al., 2013). Systemic administration of (non-selective) agonists or positive allosteric modulators of GABA_{B} receptors (GABA_{B1a/b} knock-out) impairs extinction learning (Jacobson et al., 2006), pharmacological GABA_{B} receptor antagonism does not achieve fear reduction quicker than DCS group after 1-month follow-up (Kuriyama et al., 2011).

Taken together, the available evidence indicates that GABA signaling has a major, but spatially and temporally restricted role in the formation of fear extinction memories. While GABA that is engaged during extinction is likely to support synaptic plasticity, increasing GABAergic tone globally seems to limit the efficacy of extinction-based therapies by mechanisms outlined above. Therefore, it is unlikely that drugs broadly targeting the GABA system will be useful in this respect. Pharmacological adjuncts to exposure therapy targeting the GABA system would rather need to strike a balance between driving and maintaining relevant

| Table 4A |
| Human trials: D-cycloserine (DCS) combined with CBT. |
| Disorder | Study design | Outcome (compared to placebo control group) | Reference |
|---|---|---|---|
| Healthy volunteers | DCS or placebo 2–3 h prior extinction training | No effect on fear extinction or fear recovery in healthy volunteers | (Guastella et al., 2007) |
| Healthy volunteers | DCS, placebo, valproic acid or combination of DCS and valproic acid 1.5 h prior extinction training | Administration of DCS, valproic acid or DCS/valproic acid combination facilitates fear extinction and protects from reinstatement | (Kuriyama et al., 2011) |
| Healthy volunteers | DCS or placebo 2 h prior extinction training | No effect on extinction learning and retention in healthy volunteers | (Klumpers et al., 2012) |
| Acrophobia | DCS or placebo 1 h prior 2 sessions of virtual reality exposure 3 month follow-up | Reduced fear symptoms in DCS group at all timepoints | (Ressler et al., 2004) |
| Acrophobia | 2 sessions virtual reality exposure plus DCS or placebo after the sessions 1 month follow-up Re-evaluation of (Tart et al., 2013) | No advantage of DCS over placebo augmented VRE | (Tart et al., 2013) |
| Snake phobia | DCS or placebo 1 h prior a single exposure session | Same level of improvement with DCS, but DCS group showed greater reduction in panic symptom severity at all timepoints | (Nave et al., 2012) |
| Panic disorder | DCS or placebo 1 h prior CBT session 3–5 1 month follow-up | No benefits of DCS at all timepoints, but initial beneficial effects in severely symptomatic patients? | (Siegmund et al., 2011) |
| Panic disorder with agoraphobia | DCS or placebo 1 h prior 3 individual exposure sessions + 8 group CBT sessions 5 months follow-up | DCS group showed greater reduction in panic symptom severity at all timepoints | (Otto et al., 2010c) |
| PTSD | DCS or placebo 1 h prior 10 weekly exposure sessions | No overall enhancement of treatment effects. Higher symptom reduction in severe cases. | (de Kleine et al., 2012) |
| PTSD (combat-related) | DCS or placebo 30 min prior exposure session 2–6 weeks | Weaker symptom reduction compared to placebo group | (Litz et al., 2012) |
| PTSD | DCS or placebo 1.5 h prior 12 weekly VRE session 6 months follow-up | DCS group showed earlier and greater improvement as well as higher remission rates | (Difede et al., 2014) |
| Pediatric PTSD | DCS or placebo 12 h prior session 5–12 | No difference in symptom reduction, DCS group showed trend for faster response and better retention in 3 month follow-up | (Scheeringa & Weems, 2014) |
| OCD | 1× psychoeducation; DCS or placebo 1 h prior to 4 h exposure session (5×, individual or group) 1 month follow-up | Fewer social fear and avoidance in DCS group. Significant differences in DCS vs placebo following 3rd exposure session. | (Guastella et al., 2008) |
| OCD | DCS or placebo 1 h prior to 4 h exposure therapy (5×) 1 month follow-up | Greater overall rates of improvement and lower post-treatment severity | (Smits et al., 2013b) |
| OCD | DCS or placebo with CBT | Similar response and remission rates, DCS group improved quicker than placebo group. | (Hofmann et al., 2013b) |
| OCD | 6× faster than placebo group | Fewer social fear and avoidance in DCS group. Significant differences in DCS vs placebo following 3rd exposure session. | (Siegmund et al., 2011) |
| OCD | 2×/week DCS or placebo 2 h prior exposure and response prevention (ERP) sessions (max 10) 3 months follow-up | Faster improvements in DCS group No difference between DCS and placebo group (no further benefit). | (Kushner et al., 2007) |
| OCD | 2×/week DCS or placebo 1 h prior 10 ERP sessions 1 month follow-up Re-evaluation of Wilhelms et al., 2008 | Faster improvements in DCS group No difference between DCS and placebo group (no further benefit). Specific improvements in the first 5 sessions, 6× faster than placebo group | (Wilhelm et al., 2008) |
| Pediatric OCD | DCS or placebo 1 h prior weekly session 4–10 (psychoeducation, cognitive training and ERP) for 10 wks | Modest reduction of obsessive symptoms | (Storch et al., 2007) |
| Pediatric OCD | DCS or placebo 1 h prior session 5–9 ERP-CBT | Significant improvements in OCD severity from posttreatment to 1-month follow-up in severe and difficult-to-treat pediatric OCD | (Farrell et al., 2013) |
| Pediatric OCD | DCS or placebo immediately after each of 10 CBT (ERP) sessions 1 year follow-up | Both groups improved; no significant advantage of DCS at any timepoint | (Mataix-Cols et al., 2014) |
GABA activity without over-activating the system. The evidence so far suggests that subunit-selective or allosteric (distinct from BZD) modulation of GABA<sub>A</sub> receptor signaling may hold promise in this regard.

4.6. Cannabinoids

A growing body of literature demonstrates that the endogenous cannabinoid (eCB) system modulates neuronal excitability in stressful and fearful situations (de Bitencourt et al., 2013; Gunduz-Cinar et al., 2013a). eCB signaling has been implicated in emotional and cognitive processing of threatening stimuli, for instance during the consolidation of fear and fear extinction memories (for more detailed recent reviews see (Riebe et al., 2012; Gunduz-Cinar et al., 2013b). Upon neuronal activation, an eCB (anandamide) and 2-arachidonoylglycerol (Fig. 8) are rapidly synthesized in the postsynaptic neuron (within minutes) and released into the synaptic cleft to modulate presynaptic signaling by the activation of Gi/o-coupled CB-1 receptors (CB-2 receptors are mainly expressed on immune cells, osteoblasts, osteoclasts and hematopoietic cells but also on neuroglia).

Supporting a role of eCB signaling in the extinction of aversive memories (see Fig. 8 for overview), there are increased eCB levels in the mouse BLA following extinction training (Marsicano et al., 2002).
Conversely, PTSD patients show low brain anandamide levels (Neumeister et al., 2013). In rodents, facilitation of eCB signaling by CB1 receptor agonists or exogenous cannabinoids [cannabidiol, (Bitencourt et al., 2008)], reuptake inhibitors [AM404, (Bitencourt et al., 2008; Chhatwal et al., 2005b; Pamplona et al., 2008)] or anandamide degradation blockers [AM3506, URB 597, (Gunduz-Cinar et al., 2013b)] enhances the formation or consolidation of extinction memories. Blockade or knock-out of CB1-receptors (Cannich et al., 2004; Kamprath et al., 2006; Marsicano et al., 2002; Niyuhire et al., 2007; Pamplona et al., 2006; Plendl & Wójcik, 2010; Reich et al., 2008; Terzian et al., 2011) impairs extinction and blocks the pro-extinction effects of drugs that facilitate eCB transmission (Chhatwal et al., 2005b; Do Monte et al., 2013; Gunduz-Cinar et al., 2013b) (see Table 6). Collectively, these observations point to CB-1 receptor mediated eCB signaling as a powerful modulator of extinction.

Following up on these studies, intra-cerebral infusions of CB1-receptor agonists and antagonists have revealed a potential role of mPFC-BLA eCB signaling in extinction [see Table 6, (Do Monte et al., 2013; Ganon-Elazar & Akirav, 2013; Gunduz-Cinar et al., 2013b)]. CB1 receptor activation inhibits both presynaptic glutamate and GABA release. The release of anandamide and concomitant CB1-receptor activation induce long-term depression of inhibitory transmission (LTDi) in the BLA (Azad et al., 2004; Gunduz-Cinar et al., 2013b). In addition, a recent study demonstrated that enhanced inhibitory input via activation of CB1 receptor expressing cholecystokinin (CCK) positive interneurons selectively reduces firing of BLA fear neurons and
promotes extinction (Trouche et al., 2013). Of note is the fact that it has also been shown that one potential mechanism for the CB1 effect on extinction within the AMY is via blockade of CCK release, as an intra-AMY CCK antagonist reversed the extinction blockade of systemic CB1 antagonists, and CCK agonists block extinction (Chhatwal et al., 2009).

In summary, current preclinical data suggest that eCB signaling gates BLA function through plastic changes, including a long-lasting depression of inhibition, to modulate extinction-related AMY activity. Initial results from clinical studies in healthy volunteers suggest that exogenous cannabinoid administration strengthens extinction and protects against reinstatement [see Table 6A, (Das et al., 2013; Rabinak et al., 2013)]. Future clinical trials in extinction-impaired PTSD patients will be needed to show whether cannabinoids (or substances which increase endogenous eCB signaling, such as FAAH inhibitors) might prove useful as adjuncts to CBT-based therapy.

4.7. Neuropeptides (oxytocin, neuropeptide Y, neuropeptide S, opioids)

Various neuropeptides and their corresponding receptors are localized in brain regions mediating emotional behavior and stress response and have been considered as potential anxiolytic drug targets (Bowers et al., 2012; Ebner et al., 2009; Griebel & Holmes, 2013). While other neuropeptides such as hypocretin/Orexin (Flores et al., 2014) may also be involved in fear extinction, we will here address the role of oxytocin, neuropeptide Y, neuropeptide S and opioids in extinction.

4.7.1. Oxytocin

The oxytocin (OXT) system, strongly implicated in social cognition and behavior, is posited to play a role in anxiety disorders with impaired social functioning (Meyer-Lindenberg et al., 2011; Neumann & Landgraf, 2012). OXT is synthesized in the paraventricular (PVN) and supraoptic nuclei of the hypothalamus and processed along the axonal projections to the posterior pituitary gland. Upon neuronal activation OXT is released from secretory vesicles into the peripheral circulation as neurohypophysial hormone. The central nervous system (CNS) component of OXT is derived from dendritic vesicles in the peripheral circulation as neurohypophysial hormone. The central nervous system (CNS) component of OXT is derived from dendritic vesicles in the peripheral circulation as neurohypophysial hormone. The central nervous system (CNS) component of OXT is derived from dendritic vesicles in the peripheral circulation as neurohypophysial hormone. The central nervous system (CNS) component of OXT is derived from dendritic vesicles in the peripheral circulation as neurohypophysial hormone. The central nervous system (CNS) component of OXT is derived from dendritic vesicles in the peripheral circulation as neurohypophysial hormone.
among other regions, the AMY [especially CeL (Knobloch et al., 2012; Veinante & Freund-Mercier, 1997)], the dorsal and ventral HPC and the main monoamine nuclei – substantia nigra (dopamine), raphe nuclei (serotonin) and the locus coeruleus (noradrenaline) [see (Gimpl & Fahrenholz, 2001) for review].

Regulatory effects of OXT on stress-induced activation of the hypothalamic–pituitary–adrenal (HPA) axis, as well as direct OXT effects on the brain and specific parts of the extinction circuitry, can affect fear extinction in various ways. Intracerebroventricular (icv) infusion of OXT prior to fear conditioning does not affect fear learning but facilitates fear extinction in various ways. Intracerebroventricular (icv) infusion of OXT prior to fear conditioning does not affect fear learning but facilitates fear extinction in various ways.

Table 6
Cannabinoid signaling in fear extinction (preclinical studies).

| Drug/manipulation | Extinction training | Extinction retrieval | Longterm extinction | Route | Reference |
|-------------------|---------------------|----------------------|---------------------|-------|-----------|
| Cannabidiol       | +                   | +                    | ns                  | icv   | (Bitencourt et al., 2008) |
| Anandamide        | ns                  | +                    | ns                  | IL    | (Do Monte et al., 2013)    |
| AM404 (eCB uptake inhibitor) | +                     | +                    | ns                  | dCA1  | (de Oliveira Alvares et al., 2008) |
| AM3506 (FAAH inhibitor) | +                   | +                    | (Re-in)             | ip    | (Chhatwal et al., 2005b)   |
| WIN 55.212-2 (low-dose CB1 Ag) | +                   | +                    | ns                  | icv   | (Bitencourt et al., 2008)   |
| WIN 55.212-2 (high-dose CB1 Ag) | +                   | +                    | ns                  | IL    | (Lin et al., 2009)          |
| WIN 55.212-2 (CB1 Ag) | ns                  | +                    | ns                  | IL    | (Gunduz-Cinar et al., 2013b) |
| AM1210 (AEA hydrolysis inhibitor) | ns                  | +                    | ns                  | IL    | (Lin et al., 2009)          |
| AM3506 (FAAH inhibitor) + SR141716A (CB1 ant) | +                   | No effect            | ns                  | BLA   | (Gunduz-Cinar et al., 2013b) |
| Cannabidiol + SR141716A (CB1 ant) | No training | No effect | ns                  | IL + ip | (Do Monte et al., 2013)    |

Reduced cannabinoid signaling

| Drug/manipulation | Extinction training | Extinction retrieval | Longterm extinction | Administrationroute | Reference |
|-------------------|---------------------|----------------------|---------------------|---------------------|-----------|
| CB1 KO            | —                   | —                    | ns                  | No drug             | (Cannich et al., 2004) |
| CB1 KO            | —                   | ns                   | ns                  | No drug             | (Dubreucq et al., 2010); (Kampbrath et al., 2006); (Marsicano et al., 2002); (Plendl & Wotjak, 2010) |
| D1-CB1 KO         | —                   | ns                   | ns                  | No drug             | (Terzian et al., 2011) |
| SR141716A CB1 ant | ns                  | No effect            | ns                  | sc, ip              | (Marsicano et al., 2002); (Kampbrath et al., 2006); (Niyuhire et al., 2007); (Plendl & Wotjak, 2010) |
| ns                | —                   | ns                   | ns                  | sc, ip              | (Suzuki et al., 2004) |
| ns                | —                   | No effect            | ns                  | ip                  | (Pamplona et al., 2006, 2008) |
| ns                | +                   | No effect            | ns                  | ip                  | (Chhatwal et al., 2005b) |
| ns                | +                   | No effect            | ns                  | ip                  | (Bowers & Ressler, 2015) |
| ns                | —                   | No effect            | ns                  | ip                  | (Reich et al., 2008) |
| ns                | —                   | No effect            | ns                  | dCA1                | (de Oliveira Alvares et al., 2008) |
| ns                | —                   | No effect            | ns                  | IL                  | (Lin et al., 2009) |
| ns                | —                   | No effect            | ns                  | PL                  | (Kuhnhert et al., 2013) |

1 Drug administration following extinction training; 2 drug-free retrieval of contextual fear memory; 3 5 days; 4 single-prolonged-stress model: injection immediately after trauma.

*, Recent memory; †, remote memory; #*, enhanced immobility at start of extinction training; +, Improved; −, impaired; (+) or (−), only minor effects; ip, intraperitoneal injection; ip, sc, subcutaneous injection; ns, not studied; HPC, intra-hippocampal administration; IL, infralimbic cortex; PL, prelimbic cortex; CA1, cornu ammonis 1; Re-in, reinstatement; SR, spontaneous recovery, ag, agonist; ant, antagonist, KO, knock-out;

Table 6A
Human trials: cannabinoids combined with CBT.

| Disorder         | Study design                                                                 | Outcome (compared to placebo control group)                  | Reference |
|------------------|-------------------------------------------------------------------------------|-------------------------------------------------------------|-----------|
| Healthy volunteers | Fear conditioning paradigm, cannabidiol inhalation prior or following extinction | Reduced fear in retrieval                                   | (Das et al., 2013) |
| Healthy volunteers | Fear conditioning paradigm, tetrahydro-cannabinol prior extinction           | Protection against reinstatement                             | (Klumpers et al., 2012) |
| Healthy volunteers | Fear conditioning paradigm, Dronabinol prior extinction                        | Reduced fear in retrieval                                   | (Rabinak et al., 2013) |
later extinction acquisition and retrieval (Toth et al., 2012a). Conversely, OXT receptor antagonists impair extinction learning and retrieval when administered icv prior to fear acquisition (Toth et al., 2012a). More localized OXY infusions into the central CeA reduce fear expression by inhibiting CeM excitatory output to fear-eliciting brainstem structures (Huber et al., 2005) (Viviani et al., 2011), suggesting one possible mechanism for the effects of icv OXY on extinction.

Another potential mechanism might be OXT-induced dampening of stress-induced activation of the HPA axis (de Oliveira et al., 2012; Heinrichs et al., 2003; Quirin et al., 2011; Windle et al., 1997). As explained in more detail in Section 4.8 Glucocorticoids, glucocorticoid release during an actual learning process facilitates cognitive processing [reviewed in (Benz et al., 2010)], hence the release of endogenous OXT during traumatic experiences (Zoicas et al., 2012) may interfere with the consolidation of fear memories. Along these lines, OXT-mediated reduction of HPA axis activity and subsequent lower glucocorticoid levels could interfere with fear acquisition, resulting in a weaker fear memory that is more susceptible to extinction prior to training. In this context, OXT administration either icv or intra BLA prior to extinction training disrupts extinction learning ([Toth et al., 2012a] (Lahoud & Maroun, 2013) see Table 7 for summary). OXT infusions into the dorsolateral septum prior to extinction training have the opposite effect and facilitate extinction learning and retrieval in a social fear conditioning paradigm that may recruit a somewhat different neural circuitry than non-social forms of fear (Zoicas et al., 2012).

Considering that systemically applied neuropeptides do not cross the blood–brain-barrier, clinical use of neuropeptides would require a direct pathway to the human brain. In the case of OXT, this may be feasible via intranasal administration (Born et al., 2002). Intranasal OXT administration has been demonstrated to reduce stress-induced cortisol release in humans (de Oliveira et al., 2012; Heinrichs et al., 2003; Quirin et al., 2011), and to attenuate AMY hyperactivity as well as AMY-hindbrain coupling in anxiety patients ([Kirsch et al., 2005; Labuschagne et al., 2010]). In a recent study investigating the effects of intranasal OXT treatment on fear extinction in healthy volunteers, OXT increased initial fear expression during extinction training, but this effect was gone by the end of training and there was a lower level of fear in the OXT treated group in extinction retrieval (Acheson et al., 2013). These data raise the prospect of OXT-augmented CBT effectiveness.

4.7.2. Neuropeptide Y

Neuropeptide Y (NPY) contains 36 amino acids and is the most widely expressed neuropeptide in the mammalian brain. There are particularly high concentrations of NPY in the limbic system, including in the AMY, the HPC and the periaqueductal gray. Six G-protein coupled NPY receptors (Y1–6) have been identified, among which Y1, Y2, Y4, and Y5 mediate central effects. Y1 mRNA is present at a high level in brain areas involved in memory processing, namely the AMY, HPC, thalamus, hypothalamus, and the cerebral cortex. The AMY, HPC, and hypothalamus are also rich in Y2 receptors, while Y5 receptors can be detected in HPC, cingulate cortex and thalamic and hypothalamic nuclei (Parker & Herzog, 1999) [reviewed in (Holmes et al., 2003)].

Icv administration of NPY facilitates extinction learning (Gutman et al., 2008), while genetic deletion of NPY disrupts extinction acquisition and subsequent retrieval (Verma et al., 2012). Extinction deficits caused by NPY knock-out are rescued by AAV-mediated NPY expression in the BLA (Verma et al., 2012), where NPY co-localizes with GABAergic neurons modifying inhibitory control of BLA projection neurons (McDonald & Pearson, 1989). To elucidate which NPY receptors mediate the BLA-associated effects on extinction, the consequences of gene deletion via either deletion of Y1 or Y2 receptors or double Y1/Y2 deletion have been examined. Deletion of Y1 receptors impaired extinction learning, while Y2 receptor deletion had no effect (Verma et al., 2012). Underscoring the importance to extinction of Y1 receptors in the BLA, local infusion of the Y1 agonist Leu31Pro34-NPY prior to extinction training led to stronger extinction retrieval (Gutman et al., 2008; Lach & de Lima, 2013). Further studies are required to further delineate the underlying mechanisms of Y1-mediated facilitation of extinction, and to investigate the potential clinical use of NPY and Y1 receptor agonism as adjunctive therapy to CBT.

Table 7
Neuropeptide signaling in fear extinction (preclinical studies).

| Neuropeptides | Drug/administration | Extinction training | Extinction retrieval | Longterm extinction | Route | Reference |
|---------------|---------------------|---------------------|---------------------|---------------------|-------|-----------|
| Oxytocin      | −                   | +#                  | −                   | ns                  | icv   | (Toth et al., 2012a) |
|               | +                   | +                   | −                   | ns                  | BLA   | (Zoicas et al., 2012) |
|               | −                   | −                   | −                   | ns                  | icv   | (Eskandarian et al., 2013) |
| Chronic oxytocin (30d) | ns | No effect + | ns | in | (Bales et al., 2014) |
| NPY           | +                   | +                   | +                   | ns                  | icv   | (Gutman et al., 2008) |
| NPY Y1 KO     | −                   | −                   | −                   | ns                  | No drug | (Verma et al., 2012) |
| NPY Y2 KO     | No effect           | No effect           | No effect           | No drug            | No drug | (Verma et al., 2012) |
| NPY Y1/Y2 KO  | −                   | −                   | −                   | ns                  | No drug | (Verma et al., 2012) |
| Leu31-Pro-NPY (Y1 ag) | ns | +           | ns                 | BLA                | (Lach & de Lima, 2013) |
| BIBO 3304 (Y1 ant) | +       | No effect         | No effect         | No effect | BLA | (Gutman et al., 2008) |
| NPS           | −                   | −                   | −                   | −                   | −      | (Jungling et al., 2008) |
|               | +                   | +                   | +                   | +                   | +      | (Jungling et al., 2008) |
| SHA68 (NPS-R ant) | −       | −                   | −                   | −                   | −      | (Jungling et al., 2008) |
| Naloxone      | ns                  | No effect           | No effect           | NS                 | NS     | (McNally & Westbrook, 2003) |
|               | −                   | −                   | −                   | −                   | −      | (McNally & Westbrook, 2003) |
| CTAP (MOR ant) | −                   | −                   | −                   | −                   | −      | (Parsons et al., 2010) |
| Dynorphin KO  | −                   | −                   | −                   | −                   | −      | (Parsons et al., 2010) |
| nor-BNI (KOR ant) | ns | −                   | −                   | −                   | −      | (McNally et al., 2005) |

1 Drug administration following extinction training; 2 social fear conditioning; 3 animals show accelerated fear learning; 4 administration 2h prior extinction training; 5 single prolonged stress model; 6 BTBR mouse autism model and C57BL/6j.

** Enhanced fear expression at the beginning of extinction training. + reduced fear expression at the beginning of extinction training. ** Improved; − impaired; (+) or (−), only minor effects; ip, intraperitoneal injection; icv, intra-cerebroventricular injection; ns, not studied; HPC, hippocampal administration; BLA, intra-basolateral amygdala administration; DLS, dorsolateral septum; vIPAG, ventrolateral periaqueductal gray; Ren-A, fear renewal in conditioning context; ag, agonist; ant, antagonist, KO, knock-out;
4.7.3. Neuropeptide S

Neuropeptide S (NPS) is synthesized in neurons located adjacent to the locus coeruleus (Xu et al., 2007) and binds to G-protein coupled NPS receptors expressed in the BLA among other areas. Intra-BLA infusion of exogenous NPS accelerates extinction learning, while BLA NPS receptor antagonism (via SHA68) impairs extinction learning and retrieval and increases fear renewal (Chauveau et al., 2012; Jungling et al., 2008) (Table 7). NPS treatment also rescues deficient extinction learning in hyperanxious rats (Slattery et al., in press), exhibiting aberrant cortico-AMY activation (Muigg et al., 2008). The activation of NPS receptors in the BLA increases excitatory input to GABAergic ITCs, thus increasing feed-forward inhibition to CeM neurons and reducing fear expression (Meis et al., 2008; Meis et al., 2011; Pape et al., 2010). Exogenous NPS administration is also associated with enhanced DA release in the mPFC (Si et al., 2010), suggesting another mechanism for strengthening extinction (see Section 4.2 Dopaminergic system).

Underscoring the translational relevance of NPS, healthy human volunteers carrying the NPS receptor polymorphism rs324981, which potentiates NPS efficacy at the receptor, showed enhanced extinction in a virtual reality fear potentiated startle paradigm (Glotzbach-Schoon et al., 2013). Additional studies are required to assess whether increasing NPS signaling improves extinction in anxiety patients. This would be aided by the development of small molecule NPS receptor agonists.

4.7.4. Opioids

The opioid peptide family comprises different members including endorphins, enkephalins, dynorphins and hemorphins. The 3 classical opioid receptors [μ (MOR), κ (KOR) and δ (DOR)] show heterogeneous expression throughout the brain and are located in extinction-relevant brain areas including the AMY, mPFC and HPC. Typically, MOR, KOR and DORs are coupled to Gi-proteins, reducing (in most cases) net neuronal excitability and reducing neurotransmitter release upon activation (reviewed in [Nutt, 2014]).

There is evidence that morphine application (intended for pain management) following an acute trauma attenuates the incidence of PTSD at later time points (Bryant et al., 2009; Holbrook et al., 2010; Melcer et al., 2014; Szczytkowski-Thomson et al., 2013). Aside from this potential PTSD preventing property of opioids, several studies have tried to determine the importance of opioids in extinction.

Systemic administration of the opioid receptor antagonist naloxone (which has highest affinity to MOR, but also inhibits KOR and DOR) impairs extinction when given prior to extinction training (McNally & Westbrook, 2003; McNally et al., 2004) but not when applied after extinction training or prior to retrieval, see Table 7). Subsequent investigations have demonstrated that extinction-impairing effects are mediated by MORs in the PAG [as shown with the selective MOR antagonist CTAP in the PAG (McNally, 2005; Parsons et al., 2010)].

The AMY ITCs express high levels of MOR (Busti et al., 2011), and lesioning of MOR-expressing neurons in the ITCs following fear extinction training and blunted functional connectivity between AMY and mPFC (Bilkei-Gorzo et al., 2012). Along these lines, genetic deletion of dynorphin or systemic antagonism of KORS in rodents impairs extinction learning and reduces neuronal activity in the BLA and mPFC (Bilkei-Gorzo et al., 2012). In contrast to these extinction-impairing effects, systemic administration of KOR antagonists protects against fear renewal (Cole et al., 2011) — an effect mimicked by local infusions into the ventral HPC (Cole et al., 2013). Additional studies will help clarify these effects and their possible relevance to clinical use of opioidergic drugs in conjunction with exposure-based therapies.

4.8. Glucocorticoids

Emotionally arousing experiences activate the HPA axis and the release of glucocorticoids (GCs) into the bloodstream. GCs which include cortisol in humans, and corticosterone in rodents are lipophilic and readily cross the blood–brain-barrier to interact with glucocorticoid (GR) and mineralocorticoid receptors (MR) in the brain. GCs are widely expressed throughout the brain including the AMY, HPC and PFC (Benz et al., 2010). Early pioneering studies reported that extinction-memory formation is disrupted by adrenalectomy (Silva, 1973), as well as, more recently, by compounds that antagonize GRs, such as mifepristone (Yang et al., 2006), or that inhibit corticosterone synthesis [metapyrone, (Barrett & Gonzalez-Lima, 2004; Blundell et al., 2011; Harrison et al., 1990; Yang et al., 2006, 2007]) (see Table 8). Conversely, systemic administration of corticosterone (Blundell et al., 2011) or the GR agonists dexamethasone or RU28362 (Ninomiya et al., 2010; Yang et al., 2006, 2007) facilitate the consolidation of aversive and appetitive extinction memories (see Table 8).

How can stress hormones promote the formation of extinction memories? Cytosolic GRs induce genomic responses by binding to glucocorticoid responsive elements within promoter regions of GC-responsive genes, and act as transcription factors inducing the expression of learning-related genes. In addition, membrane-bound GRs can induce non-genomic actions of relevance to extinction, including the synthesis of eCBs (see Section 4.6 Cannabinoids) from membrane phospholipids (Di et al., 2003; Hill et al., 2005, 2011). Further supporting a link between GRs and eCBs, and possibly of importance in GC-augmented extinction consolidation, adrenalnectomy leads to reduced CB-1 receptor expression (Mailleux & Vanderhaeghen, 1993) and GC-mediated memory consolidation is prevented by CB-1 receptor antagonists (Campolongo et al., 2009).

GRs can also induce indirect learning-relevant genomic actions via activation of intracellular signaling cascades such as the ERK/MAPK pathway (reviewed in Reul, 2014) in cooperation with other neurotransmitter systems or ion channels including β-adrenoceptors (Roozendaal et al., 2008), and L-type calcium channels (Karst et al., 2002). These interrelated effects are functionally relevant, given that the infusion of β-adrenoceptor antagonists into the BLA blocks the memory-facilitating effects of systemically administered glucocorticoids (Quirarte et al., 1997; Roozendaal et al., 2002).

Another important functional interaction occurs between GCs and the glutamate system. GCs potentiate glutamatergic signaling via genomic and non-genomic mechanisms, by (i) increasing the readily releasable pool of presynaptic glutamate vesicles, and (ii) enhancing NMDA and AMPA receptor surface expression through increased receptor trafficking (reviewed in Popoli et al., 2012). The concomitant activation of GC and NMDA receptors, which can occur during emotionally arousing experiences, induces downstream nuclear mechanisms that result in histone modifications (reviewed in Reul, 2014), an epigenetic mechanism implicated in extinction (see Section 4.10: Epigenetics for details). Underscoring the potential importance of emotional arousal and thus of GC release in this process, pretreatment with the anxiolytic BZD lorazepam (resulting in reduced arousal in response to the stressor) blocked epigenetic modifications on Histone H3, while application of the anxiogenic BZD inverse agonist FG7142 (Singewald et al., 2003)
induced transcription facilitating acetylation and phosphorylation on histone tails (Papadopoulos et al., 2011). These epigenetic effects could contribute to the aforementioned extinction impairing effects of BZDs (see Section 4.5: γ-Aminobutyric acid (GABA) and Table 5).

In contrast to the effects of acute GC elevations, chronic high levels of corticosterone reduce cell-surface NMDA and AMPA receptor expression (Gourley et al., 2009). This loss of critical plasticity mechanisms might be one explanation as to why anxiety patients with a history of repeated traumatic events, such as combat veterans, show greater resistance to treatment. However, a considerable proportion of PTSD patients have reduced cortisol levels (Yehuda, 2004) and small case resistance to treatment. However, a considerable proportion of PTSD patients have reduced cortisol levels (Yehuda, 2004) and small case studies suggest that there are beneficial effects of CBT and adjunctive cortisol administration in PTSD patients (Yehuda et al., 2010). A number of larger studies are under way to extend this work (NCT01108146, NCT00751855, NCT01525680). In addition, it has been found that cortisol-augmented CBT has efficacy in acrophobia (de Quervain et al., 2011), arachnophobia (Soravia et al., 2006, 2014) and social phobia ([Soravia et al., 2006] see Table 8A for a summary). Whether cortisol augmented CBT for non-phobic anxiety disorders, including also GAD, facilitates fear inhibition is currently being investigated in ongoing clinical studies (see Cain et al., 2012). Furthermore, future studies may implement more selective GC agonists than cortisol (which is also acting on MRs) to avoid non-specific side effects.

4.9. Neurotrophins and miscellaneous targets

4.9.1. Fibroblast growth factor-2

Fibroblast growth factor-2 (FGF2) is a multi-functional growth factor involved in brain development and learning-related molecular signaling cascades (reviewed in Graham & Richardson, 2011a). FGF2 signaling is associated with glutamate-mediated synaptic plasticity (Numakawa et al., 2002), L-type voltage gated calcium channel expression and activation (Shitaka et al., 1996) and phosphorylation of both MAPK (Abe & Saito, 2000) and CREB (Sung et al., 2001). FGF2 also promotes LTP in the HPC (Terlau & Seifert, 1990). Hence, FGF2 interacts with the molecular tools required for the formation and consolidation of extinction memories. FGF receptors are tyrosine kinase receptors expressed widely throughout the brain, including in areas within the extinction circuitry such as the HPC and the CeA, and FGF2 expression in the HPC and the mPFC is induced under stress (Molteni et al., 2001), thus suggesting that emotionally arousing situations requiring new learning generate increased FGF2 signaling which supports the formation of emotional memories.

FGF2 has been shown to cross the blood–brain barrier (Deguchi et al., 2000) and pioneering work demonstrates that systemic administration of FGF2 prior to or following extinction training facilitates the consolidation of extinction memories (Graham & Richardson, 2009, 2010). Local infusion into the BLA replicates the extinction-facilitating

### Table 8A

**Human trials: glucocorticoids combined with CBT.**

| Disorder                        | Study design                                                                 | Outcome (compared to placebo control group)                                                                 | Reference                      |
|---------------------------------|-----------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------|--------------------------------|
| Social phobia                   | Cortisone or placebo administered orally 1 h before a socio-evaluative stressor | Reduced self-reported fear during anticipation and exposure                                            | Soravia et al., 2006          |
| Spider phobia                   | Cortisone or placebo administered orally 1 h before exposure to a spider photograph in 6 sessions distributed over 2 weeks | Progressive reduction of stimulus-induced fear. Reduction of fear was maintained also 2 days following session ending (OFF-drug) | Soravia et al., 2006          |
| Spider phobia                   | Cortisone or placebo administered orally 1 h before 2 exposure therapy sessions Follow-up after 1 month | Significant decrease in phobic symptoms assessed with the Fear of Spiders Questionnaire                  | Soravia et al., 2014          |
| Acrophobia                      | Cortisone or placebo was administered orally 1 h before virtual reality exposure to heights Follow-up after 1 month | Cortisone-treated patients reported significantly less anxiety during exposure to living spiders at follow-up | de Quervain et al., 2011      |
| PTSD (combat-related)           | Memory reactivation task followed by intravenous administration of cortisol or saline Follow-up after 1 month | Reduced PTSD symptomatic in cortisol-treated patients No differences between cortisol and saline treated patients (initial improvement was lost after 1 month) | Suris et al., 2010            |
| PTSD                            | 10 weekly sessions of prolonged exposure therapy, cortisol or placebo 30min prior session 3–10 | Accelerated and greater decline in PTSD symptoms Caveat — includes only 2 patients (1 cortisol/1 placebo) | Yehuda et al., 2010           |
Neurotrophins and miscellaneous targets in fear extinction (preclinical studies).

| Neurotrophins | Extinction learning | Extinction retrieval | Long-term extinction | Route | Reference |
|---------------|---------------------|---------------------|----------------------|-------|-----------|
| FGFR2         | (+)                  | +                   | ns                   | sc    | (Graham & Richardson, 2009) |
|               | ns                   | +                   | ns                   | sc    | (Graham & Richardson, 2009, 2010) |
| BDNF          | No training          | +                   | + (Ren-A)            | BLA   | (Graham & Richardson, 2011b) |
| 7.8-Dihydroxyflavone | (+)                | No effect            | (+) (Ren-A)          | PL    | (Peters et al., 2010); (Rosas-Vidal et al., 2014) |
| Lentiviral transfected dominant negative form of TrkB | No effect           | ns                   | + (Ren-A)            | BLA   | (Chhatwal et al., 2006) |
| BDNF KD in HPC | −                   | ns                   | No drug              |       | (Heldt et al., 2007) |
| Val66Met BDNF SNP | −                  | ns                   | No drug              |       | (Soliman et al., 2010) |
| BDNF +/− KO   | −                   | ns                   | No drug              |       | (Psotta et al., 2013) |
| BDNF antibody | −                   | ns                   | IL                   |       | (Rosas-Vidal et al., 2014) |
| BDNF KO in forebrain | ns               | No effect#           | ns                   | PL    | (Choi et al., 2010) |
| Methylene blue, nitric oxide, histamine, and LTCCs | ns                  | +                   | ns                   | ip    | (Gonzalez-Lima & Bruchey, 2004) |
| Methylene blue | ns                  | ns                  | + (+) (Ren-A)        | ip    | (Wrubel et al., 2007) |
| Histamine     | ns                   | +                   | ns                   | CA1   | (Bonini et al., 2011) |
| Dimaprit (H2 ag) | ns                | +                   | ns                   | CA1   | (Bonini et al., 2011) |
| Ranitidine (H2 ant) | −                  | −                   | −                   | −     | (Fiorenza et al., 2012) |
| SKF9188 (histamine methyl-transferase inhibitor) | ns                  | +                   | ns                   | −     | (Fiorenza et al., 2012) |
| ns            | +                   | ns                   | BLA                  | −     | (Fiorenza et al., 2012) |
| ns            | −                   | −                   | PFC                  | −     | (Fiorenza et al., 2012) |
| ns            | −                   | −                   | −                   | −     | (Fiorenza et al., 2012) |
| L-NAME (nNOS inhibitor) | −                  | −                   | −                   | ip    | (Luo et al., 2014) |
| CamKII-Cre Cav1.2 KO | No effect          | ns                   | No drug              |       | (McKinney et al., 2008) |
| Cav1.3 KO     | No effect           | ns                   | No drug              |       | (Busquet et al., 2008) |
| Nifedipine (LTCC ant) | No effect          | ns                   | No drug              |       | (McKinney et al., 2006) |
| Verapamil (VGCC ant) | −                  | −                   | −                   | −     | (Cain et al., 2002) |
|               | No effect           | ns                   | icv                  | BLA   | (Busquet et al., 2008) |
|               | No effect           | −                   | −                   | −     | (Davis & Bauer, 2012) |
|               | No effect           | −                   | HPC                  | −     | (de Carvalho Myskow et al., 2014) |
|               | No effect           | −                   | −                   | −     | (Davis & Bauer, 2012) |

1 Drug administration following extinction training; 2 drug administration 24 h prior extinction retrieval; 3 drug administration 30 min prior extinction retrieval; 4 ABA scheme; 40 mg/kg.

*Facilitates rescue of impaired fear extinction; **facilitates extinction of remote memories; *** only in older animals (7 months), younger ones (2 months) are not affected; # reduced fear expression at the beginning of extinction training.

Effects of systemic FGFR2 (Table 9), demonstrating at least one key site of this neurotrophin’s action (Graham & Richardson, 2011b). Of significant importance, is the fact that FGFR2-augmented extinction has been associated with enhanced protection against return of fear phenomena including renewal and reinstatement ([Graham & Richardson, 2009, 2010, 2011b], see Table 9 for details) making it an interesting candidate for future clinical applications. Clinical studies investigating the potential of FGFR2 in CBT could be implemented in the near future as human trials investigating FGFR2 for angiogenesis have already shown some drugs to be safe (Laham et al., 2000).

4.9.2. Brain-derived neurotrophic factor (BDNF)

Brain-derived neurotrophic factor (BDNF) is the most abundant neurotrophin in the CNS and a key player in synaptic plasticity (reviewed in Andero & Ressler, 2012). BDNF binds with high affinity to the tropomyosin-related kinase B (TrkB) receptor and with low affinity to the p75NTR, a receptor for multiple neurotrophins (Reichardt, 2006). BDNF and TrkB are present in the HPC, AMY, PFC and the hippocampus, and are integral components of fear extinction mechanisms as blunted activity of BDNF-TrkB signaling is associated with deficient extinction retrieval in rats (Kabir et al., 2013). Conversely, increased levels of BDNF mRNA are observed in the BLA (Chhatwal et al., 2006), mPFC (Bredy et al., 2007) following successful fear extinction. The potential utility of BDNF-TrkB signaling as cognitive enhancing target is underscored by the findings that stimulating TrkB signaling by administering the TrkB agonist 7,8-dihydroxyflavone (DHF) facilitates LTP induction in the AMY (Li et al., 2011) and can augment fear extinction (Peters et al., 2010; Rosas-Vidal et al., 2014).

Haploinsufficiency of BDNF leads to deficits in extinction learning and retrieval (Psotta et al., 2013), while inhibition of the BDNF-evoked signaling cascade via transfection of a dominant negative form of the TrkB receptor specifically disrupts extinction consolidation in a fear potentiated startle paradigm (Chhatwal et al., 2006). Conversely, systemic injection of the TrkB agonist DHF promoted extinction in a stress model of impaired fear extinction and protected against the reinstatement of fear (Andero et al., 2011) (for an overview, see Table 9). In mice and humans, a single nucleotide polymorphism (SNP) in the BDNF gene (Val66Met), causing inefficient BDNF trafficking and reduced activity-dependent BDNF secretion, is associated with poor extinction (Table 9) and abnormal fronto-AMY activity (Soliman et al., 2010). Moreover, human carriers of the BDNF Val66Met SNP show reduced sensitivity to exposure-based therapies [reviewed in McGuire et al. (2014)], underscoring the potential role of BDNF in deficient fear extinction.

Studies aimed at identifying the site of BDNF-induced effects on extinction showed that deletion of BDNF in the HPC, but not in the PL (Choi et al., 2010), impairs fear extinction learning (Heldt et al., 2007). In addition, injection of BDNF antibodies (anti-BDNF) in the IL prior to extinction training disrupts acquisition and retrieval of extinction memories while anti-BDNF infusions in the PL cortex do not affect extinction (Rosas-Vidal et al., 2014) (Table 9). These findings indicate that BDNF
signaling in the HPC and the IL support the formation and maintenance of fear extinction memories. In fact, local BDNF injections in the IL produced fear inhibition for recent as well as remote fear memories even in the absence of extinction training (Peters et al., 2010; Rosas-Vidal et al., 2014). Given the possibility that BDNF infusions into the IL could reinstate fear by using the aforementioned extinction-independent paradigm (Peters et al., 2010), BDNF seems to induce signaling cascades crucial for the formation of extinction memories rather than decreasing the stability of the original fear memory. Overall these effects are consistent with a model in which the effects of BDNF are brain-region dependent, and BDNF appears to enhance synaptic plasticity of the most robust form of learning occurring in a time- and region-dependent manner.

Various targets described in this review may mediate their effects on extinction via interactions with BDNF. For example, chronic treatment with the SSRI fluoxetine (Section 4.1: Serotonergic system) facilitates extinction and enhances BDNF levels in the AMY and the HPC (Karpova et al., 2011). Valproate, an extinction-promoting inhibitor of histone deacetylation (see Section 4.10: Epigenetics), increases BDNF mRNA in the mPFC (Bredy et al., 2007). Furthermore, TrkB activation triggers the release of eCBs (Lemtriri-Chlieh & Levine, 2010) and increases the expression of CB1 receptors (Maison et al., 2009), hence potentially contributing to extinction memory formation (see Section 4.6: Cannabinoids). Finally, BDNF Met66 knock-in mice exhibit dysfunctional NMDA receptor-dependent synaptic plasticity in addition to reduced extinction acquisition (Ninan et al., 2010) that is rescued by systemic administration of α-cycloserine (Yu et al., 2009), underscoring an additional strong interaction between BDNF and the glutamatergic system [see (Andero & Ressler, 2012) for more details].

These findings suggest that BDNF signaling may be an important common signaling pathway for various systems involved in extinction. Although poor blood–brain-barrier permeability (Wu, 2005) limits the therapeutic potential of BDNF itself, TrkB receptor agonists such as DHF (see above) or LM22A-4 (Massa et al., 2010) may hold promise for CBT-adjunctive therapy in anxiety.

4.9.4. Histamine

The tuberomammillary nucleus is the main source of histamine in the brain. Histaminergic projections innervate main monoaminergic nuclei, such as raphe nuclei and locus coeruleus (Lee et al., 2005), as well as extinction relevant brain areas, including the AMY, mPFC and hippocampus (Kohler et al., 1985). To date, 4 histamine receptor subtypes (H1–4) have been identified — H1–3 are expressed in the CNS, while H4 is involved in chemotactic processes of the immune system. Local histamine infusions into the CA1 region of the HPC have been associated with improved extinction consolidation (Bonini et al., 2011). This effect is H2 receptor mediated, since the H2 agonist dimapir facilitates extinction consolidation while the histamine H2 antagonist ranitidine impairs it, when injected into the CA1, BLA or IL following extinction training (Bonini et al., 2011; Fiorenza et al., 2012). Furthermore, both histamine and the H2 agonist dimapir facilitate extinction-induced phosphorylation of Erk1, suggesting a possible molecular pathway mediating the augmenting effects of H2 receptor agonism on extinction consolidation (Bonini et al., 2011). However, although the H1 antagonist hydroxyzine has been used in some anxiety therapies (Guaiana et al., 2010), the potential clinical use of histamine drugs in particular targeting the H2 receptor for CBT augmentation has not been investigated.

4.9.5. Voltage gated calcium channels

Voltage-gated calcium channels (VGCCs) are classified into L-, P/Q-, R-, N- and T-type VGCCs (for a review see Catterall et al., 2005). A rise in intracellular calcium via NMDA receptor or VGCC activation initiates various second messenger signaling cascades that play a role in LTP formation. Due to their neuronal expression as well as their association with gene transcription the L-type calcium channels (LTCCs) Cav1.2 and Cav1.3 (Stryiessnig et al., 2006, 2014) have attracted most attention in the fear conditioning field.

Early studies have shown that systemic injections of LTCC blockers, e.g. nifedipine (that inhibits Cav1.2 and Cav1.3 channels), impair fear extinction acquisition (Cain et al., 2002). However, whole-brain gene knockout of Cav1.3 channels (Busquet et al., 2008; McKinney & Murphy, 2006) as well as conditional knockout of Cav1.2 on CaMKII-expressing principal neurons (McKinney et al., 2008) failed to replicate deleterious effects of LTCC blockers on extinction training. Pre-training ivc infusion of nifedipine has revealed no effect on extinction learning (Busquet et al., 2008), suggesting that the extinction-imparing effects of systemic nifedipine may be due to peripheral actions (Waltereit et al., 2008).

However, pre-training intra-BLA or intra-HPC infusion of nifedipine, or of another LTCC blocker, verapamil, impairs extinction retrieval 24h later (Davis & Bauer, 2012; de Carvalho Myskiw et al., 2014), indicating a role for LTCCs within these regions in the consolidation of extinction. This effect could be related to Cav1.3 mediation of synaptic plasticity and neuronal excitability in the AMY (McKinney et al., 2009) or to altered neuronal firing in the CA1 region of the HPC (Gamelli et al., 2011). However, the effect on extinction consolidation of selective Cav1.3 activation has not been assessed so far.

While activation of Cav1.2 channels produces severe neurological effects and germline gain of Cav1.3 function is deleterious (Scholl et al., 2013), selective Cav1.3 activation in adulthood has been shown to be tolerated at least in mice (Hetzenauer et al., 2006; Sinnegger-Brauns et al., 2004). Hence, short term use of such compounds, as would typically be required in combination with exposure therapy, should be possible. Taken together, these preclinical findings raise the prospect of targeting LTCCs, and Cav1.3 in particular, for the treatment of anxiety.
4.10. Epigenetics

The formation of extinction memories requires an intricate regulatory network of signal transduction, gene transcription and translation. A key step in this process involves the coordinated transcription of specific genes coding for learning-associated transcription factors, synaptic plasticity factors, neurotransmitter receptors, cytoskeletal proteins and other cellular substrates [reviewed in (Monti, 2013; Whittle & Singewald, 2014)]. Gene transcription is in turn tightly controlled by epigenetic mechanisms (Callinan & Feinberg, 2006). Epigenetic mechanisms regulate the accessibility of deoxyribonucleic acid (DNA) to transcription-initiating machinery via certain molecular events including histone tail post-translational modifications and DNA methylation. In this section we will focus on histone acetylation and DNA methylation, with emphasis on their dynamic interactions within the chromatin environment that orchestrate the regulation of genes at the molecular level, and review accumulating evidence that modulation of extinction-relevant receptors at the synapse can initiate epigenetic programs within the nucleus and modulate extinction in a persistent and context-independent manner. We will also review data showing that drugs targeting epigenetic mechanisms may serve to strengthen extinction.

4.10.1. Histone modifications

To date, a main focus of research is on understanding the interaction between histone modifications and the strengthening of fear extinction memories. In eukaryotic cells, DNA is organized in a highly conserved structural polymer, termed chromatin. The basic building block of chromatin is the nucleosome, which consists of 146 base pairs (bp) of DNA wrapped around an octamer consisting of (dimers of) core histone proteins (H2A, H2B, H3, and H4) held together by a H1 linker protein (Kornberg & Lorch, 1999). Histones contain a globular domain with a covalently modifiable N-terminal tail consisting of lysine and/or arginine residues. Histone modifications constitute a signal that is read alone or, in combination with other modifications and interactions with neighboring histones—a ‘histone code’ (Kouzarides, 2007). At least 9 different posttranslational modifications, including acetylation, phosphorylation, methylation, biotinylation, SUMOylation, ADP ribosylation, and ubiquitination, influence chromatin condensation, resulting in gene transcription by coordinating protein and enzyme complex accessibility to DNA (Ruthenberg et al., 2007). Here, we focus primarily on the most well-studied histone modification in extinction—acetylation of histone proteins.

Histone acetylation on lysine (K) residues leads to transcriptional activation by neutralizing the positive charge of the K ε-amino group of histone tails, which in turn relaxes chromatin by reducing the electrostatic bonding of histone with negatively charged DNA (Grayson et al., 2010). The enzymes involved in regulating this process are 1) histone acetyltransferases (HATs), also known as lysine acetyltransferases (KATs) (Allis et al., 2007) or ‘writers’ (Monti, 2013) and 2) histone deacetylases (HDACs), also known as lysine deacetylases (KDACs) (Choudhary et al., 2009) or ‘erasers’ (Monti, 2013). HATs acetylate K residues to promote gene transcription, whereas HDACs, which are present in co-repressor complexes, remove K residues to impair gene transcription (see Fig. 9) [for greater detail, see (Fischer et al., 2010)].

There is some evidence that extinction is associated with alterations in HAT activity and that drugs increasing HAT activity could constitute a novel approach to augmenting extinction (Fig. 9). Following extinction, there is increased expression of the HAT p300/CBP-associated factor (PCAF) in the rodent IL, and intra-IL infusion of the PCAF activator, SPV106, facilitates extinction and protects against fear renewal (Wei et al., 2012). However, inhibiting, rather than activating, the activity of another HAT, p300, in the IL also strengthens extinction (Marek et al., 2011). One possible explanation of this finding is that rather than strengthening extinction, p300 may bolster fear memory reconsolidation via transcription of genes including nuclear factor ε B (NF-ε B) (Furia et al., 2002; Perkins et al., 1997) and yin-yang 1 (YY1) (Yao et al., 2001), known to be involved in reconsolidation (Gao et al., 2010; Lubin & Sweet, 2007). These two studies that demonstrate how targeting HATs affects fear in different ways indicate the need for additional work to determine the optimal approach to facilitate extinction with HAT modulators.

There is an increasing amount of evidence to suggest that histone acetylation is an important mechanism promoting successful fear extinction (reviewed in (Whittle & Singewald, 2014)) [Fig. 9]. An important early study found that histone extinction is associated with increased acetylation of histone H4 in the promoter region of bdnf exon IV (Bredy et al., 2007). Subsequent work showed that administration of various HDAC inhibitors including TSA (trichostatin A), sodium butyrate, entinostat (MS-275), vorinostat (SAHA, suberoylanilide hydroxamic acid), VPA (valproic acid) and CI-944 can augment fear extinction (Table 10). The potential clinical utility of using HDAC inhibitors has been further strengthened by showing that CI-944, MS-275 and SAHA rescue extinction deficits in various rodent models (Griff et al., 2014; Hait et al., 2014; Matsumoto et al., 2013; Whittle et al., 2013) (see Table 10).

An additional attribute of HDAC inhibitors is their ability to extinguish remote (aged) fear memories that are normally resistant to extinction. Extinction of recent fear memories (1 day after conditioning), using a memory update-consolidation paradigm, increases HDAC2 nitrosylation (Griff et al., 2014), which facilitates the dissociation of HDAC2 from chromatin (Nott et al., 2008), leading to histone hyper-acetylation and HPC expression of extinction-relevant genes including c-Fos. By contrast, nitrosylation of HDAC2 and c-Fos expression do not increase after extinction training of remote (1 month post conditioning) fear memories (Griff et al., 2014) [Fig. 9]. Systemic administration of the HDAC inhibitor CI-994 was able to rescue remote extinction and the associated gene expression deficits. Rescue of impaired remote extinction has been observed with dietary zinc restriction, which inhibits HDAC in addition to affecting other systems (Holmes & Singewald, 2013; Whittle et al., 2010). Collectively, these two studies support the potential clinical utility of HDAC inhibitors as adjuncts to CBT treatment of fearful memories that started long after the exposure to trauma.

Further supporting a role of HDAC activity in the modulation of fear extinction mechanisms is the observation that reduced HDAC activity is associated with improvements in fear extinction (Hait et al., 2014). At present 18 different mammalian HDAC isoforms have been identified, and they are divided into four classes (Classes I–IV). Targeting individual HDAC isoforms is an important aim of drug development (Grayson et al., 2010). The majority of currently available HDAC inhibitors target multiple isoforms of the HDAC family (e.g. classes I, II, and IV), while some of the HDAC inhibitors that have proved to be successful in augmenting fear extinction (Table 10) exhibit preferential selectivity towards class-I HDACs (encompassing HDAC1, HDAC2, HDAC3 and HDAC8) (Haggarty & Tsai, 2011).

Studies using gene knockdown or over-expression suggest that targeting specific HDAC isoforms could indeed be a promising approach to modulating extinction [reviewed in (Whittle & Singewald, 2014)]. This is exemplified by the recent finding that gene silencing of HDAC2, but not HDAC1, in forebrain neurons promotes extinction (Morris et al., 2013). However, the effect of specifically inhibiting other HDAC isoforms, including HDAC3 [recently shown to facilitate the extinction of drug-seeking memories (Malvaez et al., 2013)], remains to be tested.

It is likely that the extinction-facilitating effects of HDAC inhibitors are due to activation of extinction-related gene transcription programs. The HDAC inhibitors SAHA and VPA increase, respectively histone acetylation in the promoter region of grm2b (NMDA receptor subunit 2B) (Fujita et al., 2012), and histone H4 acetylation in promoter IV of bdnf, leading to increased BDNF exon IV mRNA expression (Bredy et al., 2007). In addition, the HDAC inhibitor CI-994 increases histone H3 acetylation in the promoter region of plasticity associated genes including ifg2, ndy6, c-Fos, npas4, and arc (Griff et al., 2014). Indeed, a range of
neurotransmitter/neuromodulator systems that are known to mediate extinction, induce transcription of relevant genes via enhancement of histone acetylation, including histone H3 acetylation in the promoter region of bdnf, camk2a, creb and the NMDA receptor subunit genes grin2a and grin2b (Shibasaki et al., 2011; Tian et al., 2009, 2010; Tsankova et al., 2006). Concerning a possible convergent downstream mechanism for these effects, serotonin via activation of PKA signaling pathways (Guan et al., 2002), L-type calcium channel activation (Dolmetsch et al., 2001) and BDNF/TrkB activation (Correa et al., 2012) all activate the MAPK/ERK signaling pathway, which is crucial for the expression of gene transcription.

Table 10
Studies showing that HDAC inhibitors augment exposure-based fear extinction and rescue extinction learning deficits.

| Drug/manipulat | Extinction training | Extinction retrieval | Longterm extinction | Route | Reference |
|---------------|---------------------|----------------------|----------------------|-------|-----------|
| HDAC1 KO (HPC) | ns                  | −*                   | ns                   | No drug | (Bahari-Javan et al., 2012) |
| HDAC2 KO in forebrain CamKII neurons | ns                  | +**                  | ns                   | No drug | (Morris et al., 2013) |
| Vorinostat/SAHA | ns                  | +                    | ns                   | ip     | (Fujita et al., 2012) |
|               | ns                  | +                    | ns                   | ip     | (Matsumoto et al., 2013) |
|               | ns                  | +                    | ns                   | ip     | (Hait et al., 2014) |
| TSA (trichostatin A) | ns                  | +                    | + (SR)               | ip, HPC | (Stafford et al., 2012) |
| Sodium butyrate | ns                  | +                    | + (SR)               | ip     | (Itzhak et al., 2012) |
|               | ns                  | +                    | ns                   | ip     | (Lattal et al., 2007) |
|               | ns                  | +                    | ns                   | ip, HPC | (Lattal et al., 2007) |
| Valproic acid  | −                   | +                    | ns                   | ip     | (Bredy et al., 2007) |
|               | −                   | +                    | + (Ren-A)            | ip     | (Bredy & Barad, 2008) |
| ME-275 (entinostat) | −                   | +**                  | + (SR, Ren-N)        | ip     | (Whittle et al., 2013) |
| CI-994         | ns                  | +**                  | + (SR)               | ip     | (Graff et al., 2014) |
| FTY720 (fingolimod) | −                   | +**                  | ns                   | ip     | (Hait et al., 2014) |

* Partial extinction training: reduction of fear during the extinction training session was not to pre-conditioning levels; ** facilitates rescue of impaired fear extinction; *** facilitate remote memories combined with memory re-consolidation update paradigm; † extinction protocol based on a re-consolidation/update paradigm
+ , Improved; − , impaired; ip, intraperitoneal injection; ns, not studied; HPC, intra-hippocampal administration; Ren-A, Fear renewal in the conditioning context; Ren-N, Fear renewal in a novel context; SR, spontaneous recovery; † reduced freezing levels at the beginning of extinction training
for fear extinction (Pape & Pare, 2010) (see also Fig. 3). MAPK/ERK signaling to the nucleus could increase histone acetylation by enhancing activity of the HAT CREB-binding protein (CBP) (Chrivita et al., 1993).

4.10.2. DNA methylation

DNA methylation is a crucial mechanism for controlling chromatin remodeling in the adult mammalian nervous system (Levenson et al., 2006; Nelson et al., 2008) and can occur at multiple sites within a gene. DNA methylation is a direct chemical modification of a cytosine side-chain that adds a methyl (–CH3) group through a covalent bond, catalyzed by a class of enzymes known as DNA methyltransferases (DNMTs) (Okano et al., 1998). The DNMTs transfer methyl groups to cytosine residues within a continuous stretch of DNA, specifically at the 5′-position of the pyrimidine ring (5-methylcytosine) (Chen et al., 1991). Not all cytosines can be methylated; typically, cytosines are followed by a guanine in order to be methylated (Day & Sweatt, 2011) (Fig. 9). However, given the recent finding that methylation can occur outside CpG islands and in CpH (where H = adenine/cytosine/thymine) islands and that these CpH islands can repress (Guo et al., 2014) substantially expands and adds further complexity to how methylation can regulate gene transcription in the central nervous system. Furthermore, this finding raises the open question of whether CpH methylation is a molecular mechanism regulating fear extinction.

In a pioneering study, linking DNA methylation and extinction, extinction-resistant female mice were found to exhibit increased DNA methylation of Bdnf exon IV and a concomitant decrease in BDNF mRNA expression in the PFC (Baker-Andresen et al., 2013). The fact that DNA methylation is reversible suggests that enhancement of gene transcription by DNA demethylation could represent a novel mechanism for strengthening extinction that has the potential to be pharmacologically exploited. Along these lines, the ten-twelve translocation (Tet) family of methylcytosine dioxygenases, which includes Tet1, Tet2, and Tet3 enzymes, catalyze oxidation of 5-methylcytosine to 5-hydroxymethylcytosine and thereby promotes DNA demethylation (Ito et al., 2011; Tahiliani et al., 2009).

Two recent studies demonstrate the importance of these Tet enzymes in extinction. Extinction leads to a genome-wide redistribution of 5-hydroxymethylcytosine and Tet3 occupancy to extinction-relevant genes including the GABA_A clustering protein, gephyrin (see Section 4.5: γ-aminobutyric acid (GABA)), and to enhanced gephyrin mRNA expression in the IL (Li et al., 2014). Furthermore, gene knockdown of Tet1 impairs extinction (Rudenko et al., 2013). While agents targeting Tet enzymes are not currently available, these emerging preclinical findings may provide the impetus to develop such compounds.

4.10.3. Non-coding RNA

In addition to histone modifications and DNA methylation, gene expression is also regulated by non-coding RNAs including micro-RNAs (miRNAs) (Spadaro & Brady, 2012). miRNAs represent classes of non-coding RNAs that mediate post-transcriptional regulation of gene expression, including genes associated with synaptic plasticity, learning and memory (Barry, 2014). Although miRNAs are usually associated with preventing mRNA translation and thereby with suppressing new protein synthesis, a specific miRNA was recently shown to augment fear extinction. Fear extinction enhanced the expression of miR–128b in the IL associated with down-regulation of plasticity-related genes including creb1 and sp1 in tandem with inhibition of the original fear memory (Lin et al., 2011). Clearly, further clarification of the role of non-coding RNAs during fear extinction is warranted, including through the development of tools by which specific non-coding RNAs can be modulated.

5. Discussion and final conclusions

In this review we aimed to summarize (druggable) systems involved in fear extinction and how pharmacological compounds have been utilized to augment fear extinction in preclinical and small-scale clinical studies. It is anticipated that these findings are paving the way for clinical use of such approaches given that extinction training is procedurally similar to exposure-based psychotherapy and the underlying fear/extinction circuitries and molecular mechanisms are well conserved across species (Milad & Quirk, 2012). As mentioned in the introduction, the main problems with current exposure-based CBT treatment of fear, anxiety and trauma-related disorders are that a significant proportion of patients either: 1) have difficulties bearing the demanding and exhausting process of this therapy, 2) do not, or not sufficiently, respond to the therapy — failing to substantially reduce their fear response, or 3) respond initially, but suffer from return of fear phenomena hampering remission and full recovery. In this review we summarize the pharmacological approaches that have been investigated and utilized to improve these shortcomings. Initial attempts to combine exposure-based CBT with pharmacotherapy included benzodiazepines that damped anxiety but also reduced the learning effects of this procedure hence provoking return of fear symptoms upon discontinuation (Hofmann, 2012; Otto et al., 2002, 2010a). A complicating issue is the observation that the interoceptive state induced by such anxiolytics form part of a “context,” and consequently may promote fear in subsequent testing in an “off drug state”, leading to state-dependent learning due to a change in interoceptive context (Maren et al., 2013).

However, there is evidence that targeting novel anxiolytic systems such as the NPS system or fibroblast growth factor-2 may elicit acute anxiolytic effects without producing sedation, state-dependency and/or interference with the formation of extinction memories. These findings are particularly exciting as they are beginning to address one of the important problems associated with CBT (i.e. intolerability to CBT exposure). However, more research is necessary to prove the utility of such approaches. As well as improving tolerability to exposure, improvements concerning onset, magnitude and duration of the therapeutic effect of exposure-based CBT are also in the main focus of drug development in this field. As outlined in this review, the augmentation of fear extinction/exposure therapy with a range of different drugs acting as cognitive enhancers to strengthen extinction memories represents an important approach to enhancing therapy efficacy. This has been made possible by revealing the neurochemical/neurobiological responses within the neural circuitry whose activity is required for fear extinction and, by extension, to rescue fear extinction deficits.

Considerable progress has been made in defining the neural basis of fear extinction and fear extinction deficits (for recent reviews see (Milad & Quirk, 2012; Holmes & Singewald, 2013; Parsons & Ressler, 2013). The steadily growing knowledge concerning where and how fear extinction is processed and how it can be pharmacologically augmented is an essential platform from which to identify promising candidates for extinction-promoting therapeutics. Indeed, by successfully translating findings from animal models, progress has been made in enhancing the outcomes of exposure-based CBT using compounds such as DCS, yohimbine and glucocorticoids as cognitive enhancers in the treatment of patients with anxiety disorders, PTSD and OCD (Hofmann & Smits, 2008; McGuire et al., 2014; Norberg et al., 2008).

As outlined in this review, an impressive number of additional extinction augmenting drugs, acting on a wide variety of pharmacological targets, have indeed been identified. Data arising from the use of animal models displaying deficient fear extinction, reflecting patients resistant to exposure therapy, indicate that different neurotransmitter/neurobiological signaling pathways are involved in different temporal phases of extinction. Hence, it is likely that different pharmacological approaches will be required to either induce fear reductions during extinction training or rescue/boost the consolidation of fear extinction (reviewed in (Holmes & Singewald, 2013). The important finding that within-session fear reduction/habituation favors drug-augmented extinction rather than facilitating reconsolidation of existing fear memories (Graham et al., 2011; Merlo et al., 2011) has also been demonstrated in clinical trials using DCS ((Hofmann, 2014) and yohimbine (Smits
et al., 2014). Moreover, multi-target approaches that simultaneously modulate the activity of diverse neurotransmitter/neurobiological systems seem to be an important option to pursue, in particular for individuals with severe extinction deficit/resistance to exposure therapy. Interestingly, in the much more well established medical fields of infectious disease, hypertension, oncology, and similar areas, targeting multiple different mechanisms of action is also the norm, not the exception and it is also considered promising in other fields of psychiatry (Millan, 2014).

One of the most important aspects of extinction-promoting drugs is their ability to bolster extinction by enhancing memory consolidation. It is likely that this extinction enhancement occurs via the enhancement of cortico-AMY synaptic plasticity through multiple mechanisms (reviewed in (Myers & Davis, 2007; Sierra-Mercado et al., 2011; Orsini & Maren, 2012; Duvari & Pare, 2014), ultimately providing lasting effects and better protection from return of fear phenomena. If translated into clinical use, these effects would support sustained symptom remission and recovery. Some of the drug targets outlined in this review (e.g. epigenetic targets including histone modification, BDNF and facilitated dopaminergic and fibroblast-growth-factor-2 signaling) and also cholinergic targets — see (Zelikowsky et al., 2013)] seem to be promising in this respect, as they promote long-term and context-independent fear inhibition when combined with fear extinction. Context-independent extinction learning (extinction generalization) offering enhanced protection against fear return phenomena such as renewal, may be supported not only by targeting specific pharmacological mechanisms, but also by behavioral approaches (see below). Fortunately, some of the promising substances in that respect are approved for human use (e.g. Valproate, L-DOPA, FG2) and thus, their extinction-augmenting capacity could be immediately examined in anxiety and trauma-focused clinical trials. Although some of the discussed agents may share important common downstream signaling pathways, it seems clear that there are a number of different pharmacological pathways to access the circuits that underlie fear extinction learning and memory (see Fig. 3), which will enable even more specific targeting of the essential extinction-promoting mechanisms in the future.

Despite this clear progress there remain major challenges for the development of medications to improve the effectiveness of exposure-based therapies, including challenges related to the design of safe, brain-penetrant molecules with limited adverse side-effects. Individual differences in treatment tolerability and efficacy, caused by genetic variation, and prior medication history are further complicating concerns. For example, the careful dissection of the brain regions mediating fear extinction including the AMY, PFC, HPC and other regions has also shown that some systems can promote both extinction-facilitating and extinction-impairing effects depending on the brain area/cellular substrate they are acting on. An example is the β-adrenoceptor, blockade of which interferes with extinction when restricted to the IL, but promotes extinction when the BLA is targeted — hence it is difficult to predict the net effect of medicating a patient with a β-adrenoceptor blocker or agonist during exposure therapy. The targeting of specific intracellular signaling cascades might be a solution for future, more selective approaches in this direction. Research in the last few years has revealed that the specific ligands can bias the signal output of G-protein coupled receptors by stabilizing the receptor structure in a distinct way to the natural ligand. By providing a means to produce functional selectivity, biased agonists and antagonists may be utilized to activate preferential, even unnatural, signaling cascades mediating the therapeutically favored downstream biological consequences [for a recent review see (Luttrell, 2014)].

Another important aspect for future clinical implementation of drug-induced facilitation of exposure therapy outcome concerns the time course of drug administration. While current drug treatment for anxiety and trauma-related disorders involves mainly chronic treatment (e.g. with SSRIs, SNRIs, TCA, MAOI) the compounds discussed in this review are (with the exception of antidepressants) recommended to be applied acutely in combination with psychotherapeutic interventions for a limited number of exposure trials. This strategy includes several advantages: First, chronic application of drugs can induce very different biological effects from those observed with acute administration. It is extremely challenging to predict how and at which stage (recall of fear memory, learning, reconsolidation, consolidation, retrieval of extinction memory) chronically applied drugs will enhance/interfere with fear extinction. For example, it is likely that facilitation of extinction and disruption of fear reconsolidation, both leading to fear reduction, depend on opposing molecular processes, such that chronic administration of the same drug may lead to very different therapeutic outcomes depending on which process (i.e. extinction or reconsolidation) it enhances (Graham et al., 2011). Second, acute administration implies a reduced side effect potential. One example is the addiction potential of some of the compounds currently examined for extinction augmentation (e.g. MDMA). Indeed, first clinical trials reported that none of the involved patients developed a substance abuse problem when MDMA was administered acutely to assist psychotherapy (Mithoefer et al., 2013).

In addition to pharmacological parameters, the efficacy and duration of exposure therapy in humans may also depend (as shown in animal studies) on manipulation of specific behavioral extinction training parameters [see also e.g. (de Carvalho, Myskiw et al., 2014; Schiller et al., 2012)] such as providing sufficient extinction training, enhancing the number of contexts for such training, trial spacing, and other approaches (reviewed in Herry et al., 2010; Fitzgerald et al., 2014a). It will be an important future line of research to test the optimal combinations between such behavioral interventions and the outlined pharmacological approaches in order to maximize treatment outcome. It is also important to note that thought needs to be given to a variety of factors concerning how to optimally utilize adjunctive and novel treatments. Such factors include consideration of monotherapy vs. add-on adjunctive treatment; use of medications in drug-naïve vs. already medicated subjects; combined use of medication explicitly with exposure/extinction psychotherapy vs. medication taken without explicit psychotherapy; timing of medication — e.g. at night to aid memory consolidation during sleep vs. immediately before or after exposure; and use of medications in countries with well-developed vs. poorly developed health systems. As the field of psychopharmacotherapy matures beyond the models that it has followed for decades, all of these considerations are important for new drug development. For example some medications may not be as efficacious in subjects with a history of antidepressant use, as has been suggested for β-cycloserine (Werner-Seidler & Richardson, 2007). Furthermore, as sleep is increasingly thought to be associated with extinction learning, targeting medication to enhance the memory consolidation process during sleep may be important (Pace-Schott et al., 2012). Finally, medications which are available readily in generic form and that do not require specialized adjunctive psychotherapy may be more readily disseminated to locations with less well-developed mental health care systems than those that are novel, or more expensive, or that may require specialized combined therapy. Taken together, these considerations remind us that identifying molecular mechanisms and novel drug approaches to fear regulation is only part of the challenge, and that the context of the dissemination of such new therapeutics is also of critical concern.

In summary, based on the evidence outlined in this review, the next few years promise to produce several novel clinical approaches to treating trauma- and anxiety-related disorders in a more efficient and persistent way that will most likely lead to enhanced symptom remission. Progress will depend on translational research approaches and a close interaction between preclinical and clinical researchers. Fig. 10 outlines how the different levels of understanding, from genetics to epigenetics to neural circuits, can inform each other as well as how they can be informed across species due to the high level of convergence of shared fear-related
processing across mammals. This research and the considerable number of compounds to choose from in the growing pharmacological tool box in this field, many of which display minimal side effects, may enable psychiatrists/psychologists to individualize drug treatment according to specific symptoms of anxiety disorders, PTSD and OCD. Such options are expected to further improve treatment outcome and to reduce the amount of psychotherapy sessions needed, or the duration or intensity that is required for long-lasting treatment success. As outlined above, evidence is beginning to shape the hypothesis that specific classes of compounds and cognitive enhancers may be superior to others in their ability to support long-term fear-inhibiting effects. There is evidence that the efficacy of different classes of drugs may be distinctly influenced by a number of factors including age, gender and other examples of individual differences. Hence, future research is warranted in order to investigate whether specific drugs used as cognitive enhancers in this field are more or less appropriate for specific individuals, conditions and treatment durations. In conclusion, cognitive enhancers present a promising option to safely augment extinction-based anxiety- and trauma therapy and to reduce treatment duration, thereby facilitating treatment coherence and outcomes.

Conflict of interest statement

Drs. Singewald, Schmuckermair, Whittle and Holmes declare no conflict of interest. Dr. Ressler is a founding member of Extinction Pharmaceuticals/Therapade Technologies. He has received no equity or income from this relationship within the last 5 years. The terms of these arrangements have been reviewed and approved by Emory University in accordance with its conflict of interest policies.

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Fig. 10. Strategy for translating research from humans to mice and back, focusing on human disorders of fear dysregulation. Rodent models of enhanced fear and impaired extinction (induced by genetic or environmental manipulations) closely resemble human symptomatology, particularly the fear dysregulation that occurs in PTSD, panic, and phobic disorders. Using these models increases the chances of identifying candidate genes for human anxiety disorders by reducing the problems of genetic heterogeneity and a variable environment. It is also important to study the involvement of these candidate genes in patients, as well as using unbiased genome-wide association studies and genome-wide epigenetic approaches to identify previously unknown genetic pathways contributing to risk in humans. Subsequently, elucidating the function of the identified gene and its epigenetic regulation using rodent models as well as the human population increases our knowledge of the neurobiology of fear disorders, which, at the same time, is necessary to improve the models. It is also necessary to use rodent models to test the ability to pharmacologically enhance extinction of fear and diminish fear expression prior to clinical test phases. Arrows demonstrate how the different levels of understanding, from genetics to epigenetics to neural circuits, can inform each other, as well as how they can be informed across species due to the high level of convergence of shared fear-related processing across mammals.

| Human Disorders of Fear Regulation | Rodent Models of Fear Regulation |
|-----------------------------------|----------------------------------|
| **PTSD, Panic Disorder, OCD, Phobias** | **Fear Conditioning, Stress, Extinction** |
| Gene – Environment Interactions | Genetic Strains & Mutant Lines with enhanced fear or impaired extinction |
| Impairment in natural recovery | Manipulate genes to understand mechanism: Knockout, knockin, siRNA, inducible viral approaches + drug targets |
| Exposure based psychotherapy | Neurobiology of Extinction Circuits: Gene-circuit interactions; optogenetics; DREADDS identification of site-selective drug targets |
| Genes which modulate fear extinction | Enhancing extinction in genetic & behavioral mouse models of extinction deficits and enhanced fear |
| Candidate Gene, GWAS, and Epigenetics | Clinical Trials of Drugs/Methods to Enhance exposure-based psychotherapy |
| Epigenetic correlates to human fear disorders | Neural Circuits governing fear extinction |
| Clinical Trials of Drugs/Methods to Enhance exposure-based psychotherapy | Rodent Model of Fear Regulation |
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