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(Article begins on next page)

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### Review

### Brain sites involved in fear memory reconsolidation and extinction of rodents

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19	Local temporary reversible treatments
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23	Prelimbic cortex
24	Entorhinal cortex

### ABSTRACT

Fear memory is a motivational system essential for organisms survival having a central role in organization of defensive behaviors to threat. In the last years there has been a growing interest on conditioned fear memory reconsolidation and extinction, two specific phases of memorization process, both induced by memory retrieval. Understanding the mechanisms underlying these two mnemonic processes may allow to work out therapeutic interventions for treatment of human fear and anxiety disorders, such as specific phobias and post-traumatic stress disorder. Based on the use of one-trial conditioning paradigms, which allow to follow the evolution of a mnemonic trace in its various phases, the present paper has attempted to reorganize the current literature relative to the rodents highlighting both the role of several brain structures in conditioned fear memory reconsolidation and extinction and the selective cellular processes involved. A crucial role seems to be play by medial prefrontal cortex, in particular by prelimbic and infralimbic cortices, and by distinct connections between them and the amygdala, hippocampus and entorhinal cortex.

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### Contents

Dea	omena $_{\overline{\Lambda}}$ introductionstructures involved in fear memory reconsolidation		00
	tructures involved in fear memory reconsolidation	••••••	00
2.1.	Amygdala		
	2.1.1. Neurotransmitter systems		00
	2.1.2. Protein kinases		00
	2.1.3. Gene expression and protein synthesis		00
2.2.	Hippocampus		
	2.2.1. Neurotransmitter systems		
	2.2.2. Protein kinases		00
	2.2.3. Gene expression and protein synthesis		00
2.3.	Cortex and other neural sites		00
. Brai	structures involved in fear memory extinction		00
3.1.	Amygdala		00
	3.1.1. Neurotransmitter systems		00
	3.1.2. Protein kinases and phosphatases		00
	3.1.3. Gene expression and protein synthesis		00
3.2.	Hippocampus		00
	3.2.1. Neurotransmitter systems		
	3.2.2. Protein kinases		00
	3.2.3. Gene expression and protein synthesis		

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## **ARTICLE IN PRESS**

E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

	3.3.	Cortex a	and other neural sites	00
		3.3.1.	Neurotransmission systems	00
		3.3.2.	Protein kinases	00
			Gene expression and protein synthesis	
4.	Discu	ssion		00
	4.1.	Neural o	rircuit underlying fear memory reconsolidation and/or extinction	00
	4.2.	Tempor	al and biochemical signatures in the neural sites involved in reconsolidation and extinction	00
	4.3.	Reconse	lidation and extinction in human anxiety disorders	00
	Ackno	owledgm	ents	00
	Refer	ences		00

### 1. Prolegomena – introduction

Fear memory is one of the most studied memories in general and 02 48 especially in rodents. It is easily and guickly learned and retained 49 for a long time even for a lifetime. For these characteristics fear 50 memory is frequently used as an experimental model to study the 51 cerebral mechanisms involved in learning and memory (Baldi and 52 Bucherelli, 2012; LeDoux, 2000). The characteristics of fear mem-53 ories acquisition and retention are amenable to investigate both 54 the brain sites involved and the phases in which they are impli-55 cated. Fear learning entails single pairing between the conditioned 56 stimulus (CS) and the unconditioned stimulus (US) which is suf-57 ficient to establish it. In rodents "one-trial" paradigms are usually 58 employed to induce aversive conditioning, such as "fear condition-59 60 ing" and "inhibitory avoidance". The "one-trial" procedures have an important feature as they allow to follow the evolution of a mem-61 ory trace in its various phases, from its origin to its disappearance, 62 and it is very useful to know the exact starting point for engram 63 formation (Ambrogi Lorenzini et al., 1998; Muller et al., 1997; 64 Sacchetti et al., 1999a). This is not possible using multi-trial condi-65 tioning paradigms that require several temporally spaced training 66 sessions. This aspect is not of secondary importance because the 67 association phase between CS and US stimuli, called "acquisition", 68 is followed by a phase defined "consolidation" during which the 69 mnemonic trace is progressively stabilized becoming increasingly 70 resistant to destruction. This strengthening allows a labile memory 71 (working memory or short-term memory) turning into a con-72 solidated memory (long-term memory) that can be stored and 73 74 retained for a very long time (Abel and Lattal, 2001; McGaugh, 2000). 75

Indeed, in the brain the life of the engram is certainly more dynamic than shown by this simplistic treatment. A wellconsolidated engram is not unchangeable. Many studies have demonstrated that a "dormant" memory trace in the brain is well protected from potential erasing, but when this trace is recalled (retrieved) it can change. The re-activation returns the engram to a labile state making it sensitive to disruption (Alberini, 2005; Nader and Hardt, 2009; Nader et al., 2000). It is necessary to make a distinction because if a new re-exposure to the CS reactivates the trace, the resulting effects depend on the re-exposure features. Short re-exposition to the CS also in the absence of the reinforcement elicits the conditioned response starting at the same time a new process of memory trace elaboration, called reconsolidation. On the contrary, prolonged or repeated re-exposures to CS alone determine a gradual weakening of the engram showing extinction (Eisenberg et al., 2003; Pedreira and Maldonado, 2003; Suzuki et al., 2004). As for the original phases of the process of engram formation (acquisition, consolidation, retrieval) reconsolidation and extinction as well can be selectively investigated employing experimental paradigms. In addition, in the processing of memory trace it is important to investigate the mechanisms that regulate it. In this regard, there is a wide scenario that involves neurotransmitters, neuromodulators, biochemical and genetic expression processes that continuously interfere with each other (Myers and Davis, 2007; 99 Quirk and Mueller, 2008; Tronson and Taylor, 2007).

To assess whether a mechanism or a brain site is involved in a 101 specific phase of the memorization process (acquisition, consolida-102 tion or retrieval of mnemonic trace), experimental techniques are 103 used that affect selectively a phase of the process without inter-104 fering with the other ones preceding or following the investigated 105 phase. For example, irreversible lesions of cerebral sites are not 106 suitable for this purpose. On the contrary, temporary (reversible) 107 inactivation allows to investigate not only the possible role of a 108 particular brain structure in mnemonic process, but also in which 109 phase or phases it is involved (Ambrogi Lorenzini et al., 1999). 110

In rodents, Pavlovian fear conditioning has been the main condi-111 tioning procedure used. In a typical experiment of fear conditioning 112 to examine fear memory, an animal is placed in a conditioning 113 apparatus and an emotionally neutral stimulus (such as a tone, 114 a light or an odor) is paired with an aversive stimulus such as a 115 mild electric footshock (the unconditioned stimulus, US). As a result 116 of this pairing, the initially neutral stimulus (being now a condi-117 tioned stimulus, CS) acquires the ability to elicit a typical behavioral 118 fear response. However, in this procedure the US is associated not 119 only with a discrete CS but also with the environment in which 120 the US is presented, i.e. the training context. Thus, the animal will 121 exhibit conditioned fear to both CS (cued fear conditioning) and 122 context (contextual fear conditioning) during the subsequent re-123 exposure to that CS or context. In this way, the same paradigm 124 allows to obtain two distinct mnemonic traces, that can be stud-125 ied separately, and to follow their temporal evolution (LeDoux, 126 2000; Sacchetti et al., 1999a, 1999b). The conditioned fear can be 127 measured by quantifying specific behavioral responses such as fear-128 potentiated startle and freezing or immobility. The former consists 129 in an increase in the amplitude of an acoustically elicited startle 130 response, whereas the latter is defined as the complete absence of 131 somatic movements except those requested for respiration (Fendt 132 and Fanselow, 1999; Sacchetti et al., 1999a, 1999b). Contextual fear 133 may also be induced by the presentation of a US alone. In addition 134 to classical fear conditioning, this form of fear memory has been 135 studied using the inhibitory avoidance paradigm. The animal learns 136 that performing a response (for example, walking from an illumi-137 nated compartment to a spontaneously preferred darkened one of 138 an apparatus or moving from a small elevated platform to a larger 139 arena) is punished with a footshock. In this case the animal learns to 140 avoid the punishment inhibiting its natural response (Tinsley et al., 141 2004). 142

As in all memories, fear memory retrieval of a consolidated 143 mnemonic trace is a dynamic process that can initiate two pro-144 cesses: reconsolidation and extinction. In the reconsolidation, 145 the retrieved fear memory transiently returns to a labile state 146 and requires a new round of consolidation to be preserved. It 147 has been proposed that reconsolidation allows the integration of 148 new information into the original mnemonic trace, thus allowing 149 memory updating, and also to strengthen or weaken it (Alberini, 150 2005; Alberini and LeDoux, 2013). Reconsolidation is not exactly 151

### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

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the fact that these sites play a central role in the fear responses 215 learning. Indeed, the amygdala was demonstrated to be critical for 216 the acquisition, consolidation and expression of cued and contex-217 tual fear conditioning. In addition, results obtained by inhibitory 218 avoidance experiments show that this neural structure is also 219 important for this form of fear memory, although in this case it 220 seems to have a modulatory role as amygdala lesions attenuate, 221 but do not block, inhibitory avoidance learning (Parent et al., 1994, 222 1995; Parent and McGaugh, 1994). 223

### 2.1. Amygdala

The involvement of the amygdala, in particular the basolateral complex (BLA), in the fear memories reconsolidation (Table 1) has been highlighted by means of tetrodotoxin (TTX) or lidocaine functional inactivation, blockers of voltage-dependent sodium channels, thus impeding the initiation and propagation of action potentials. Intra-BLA lidocaine or TTX infusion immediately after reactivation session of fear-related memory impairs both freezing response to context (Baldi et al., 2008; Bucherelli et al., 2006) and auditory CS (Sacchetti et al., 2007) and inhibitory avoidance response (Prado-Alacalà et al., 2006; Tzeng et al., 2012). However, Prado-Alacalà et al. (2006) have reported that postreactivation administration of TTX into the amygdala produced a transient amnesic effect of inhibitory avoidance which recovered with repeated retention testing. Thus, according to the Authors, "the impairment induced by post-retrieval treatment is likely due to temporary impairment of memory retrieval". It was also shown that increasing the strength of conditioning, using an US strong enough to induce generalization, the BLA role in auditory fear reconsolidation is no longer crucial (Baldi et al., 2008; Sacchetti et al., 2007). This effect may be due to other neural sites which maintain this fear memory. On this point, some Authors (Sacchetti et al., 2007) have observed that stronger auditory fear memories are affected by the combined but not independent BLA and cerebellar vermis TTX blockade.

### 2.1.1. Neurotransmitter systems

The molecular mechanisms of fear memories reconsolidation 250 are starting to be elucidated. Many studies have employed some 251 post-retrieval treatments previously used to characterize memory 252 consolidation. The results have provided evidence that the molecu-253 lar mechanisms of these two mnemonic phases are similar, but not 254 identical. Fear memory reconsolidation requires the activation of 255 several neurotransmitter systems into the BLA, such as glutamate, 256 noradrenaline and endocannabinoid. The glutamate acts through 257 two types of ionotropic receptors: NMDA (N-methyl-p-aspartate) 258 and AMPA ( $\alpha$ -amino-3-hydroxy-5-methyl-4-isoxazole proprionic 259 acid). The blockade of NMDA-receptors in the BLA before mem-260 ory reactivation blocks the beginning of reconsolidation (Mamou 261 et al., 2006), whereas the injection of a NMDA partial agonist, p-262 cycloserine (DCS), in the same neural site before the reactivation 263 session enhances fear memories (Lee et al., 2006). Further stud-264 ies reported that different subtypes of NMDA receptor within the 265 BLA may mediate memory destabilization and re-stabilization after 266 retrieval. In particular, into the BLA the NR2B subtype appears 267 to be required for auditory fear memory destabilization, whereas 268 the NR2A subtype seems to be involved in re-stabilization of this 269 fear memory (Milton et al., 2013). On the contrary, AMPA receptor 270 blockade (Mamou et al., 2006; Milton et al., 2013) does not seem 271 to have any effect on memory reconsolidation. However, a recent 272 study (Hong et al., 2013) showed that AMPA receptors may play 273 an unexpected physiological role in guiding fear memory recon-274 solidation into the BLA. In fact, the authors demonstrated that an 275 exchange from calcium impermeable AMPA (CI-AMPA) receptor 276 to calcium permeable AMPA (CP-AMPA) receptor occurs during 277

a recapitulation or a repetition of initial consolidation. In fact, the 152 time course of the two processes is different and the brain struc-153 tures and molecular processes involved may be not necessarily 154 coincident (Alberini, 2005; Bucherelli et al., 2006; Tronson and 155 Taylor, 2007). On the other hand, during retrieval session, memory 156 reactivation by means of longer re-exposition to the CS, without the 157 US presentation, triggers the extinction process that leads to pro-158 gressive reduction in the expression of conditioned fear response. 150 However, extinction does not directly modify the original fear 160 memory but leads to the formation of a new association (CS-no US) 161 that competes with the original engram, masking it. Thus, extinc-162 tion implies new learning. On the other hand, phenomena such 163 as spontaneous recovery (that is, reappearance of an extinguished 164 fear memory with the passage of time), renewal (recovery of an 165 extinguished fear memory when the CS is presented in a context 166 different from that in which extinction training took place) and 167 reinstatement (reappearance of an extinguished fear memory fol-168 lowing exposure to unsignaled US after extinction training) show 169 that the original memory is not erased but remains encoded in the 170 brain being not expressed during extinction (Baldi and Bucherelli, 171 2010; Myers and Davis, 2007; Quirk and Mueller, 2008). 172

173 The current understanding of the neural circuitry of fear memory reconsolidation and extinction is much poorer than that 174 concerning the acquisition and consolidation phases. Experimen-175 tal results indicated that these mnemonic phases are characterized 176 by both distinctive and coincident features of the anatomical and 177 molecular requirements (Alberini, 2005; Berman and Dudai, 2001; 178 Bucherelli et al., 2006; Chen et al., 2005; Izquierdo et al., 2006; Lee 179 et al., 2004; Lin et al., 2003b; Szapiro et al., 2003; Vianna et al., 2001). 180 Understanding the mechanisms of fear memory reconsolidation 181 and extinction may have clinical importance for the treatment of 182 human anxiety disorders, such as specific phobias, panic disor-183 der and post-traumatic stress disorder. In this contest treatments 184 involving reconsolidation and extinction procedures have been 185 recently used to reduce the expression of fear memory (Alberini, 186 2005; Auber et al., 2013; Davis et al., 2006; Hartley and Phelps, 187 2010; Monfils et al., 2009; Myskiw et al., 2014; Nader, 2003; Parsons 188 and Ressler, 2013; Quirk et al., 2010; Rao-Ruiz et al., 2011; Rossato 189 et al., 2010; Schiller et al., 2010; Todd et al., 2014). Thus, the iden-190 tification of both neural structures underlying the two memory 191 phases, and pharmacological agents that impair reconsolidation 192 or potentiate extinction appears to be crucial. In this context, this 193 review attempts to reorganize results in the literature aimed to 194 highlight the role of several cerebral structures in fear memory 195 196 reconsolidation and extinction in rodents, highlighting whenever possible the selective cellular processes involved. Following the 197 principles exposed above, in this review we will consider studies 198 related to one-trial aversive conditioning to investigate the role of 199 brain structures in specific phases of memorization processes. 200

### 201 2. Brain structures involved in fear memory 202 203 204 204 205 205 206 206 206 206 206 207 208<

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Experimentally, reconsolidation process of fear memory can be verified by reactivating a well consolidated mnemonic trace. For this purpose, the previously conditioned animal is subjected to a reactivation session (usually at least 24 h after training) during which the CS is briefly re-presented usually in the absence of the US. Immediately after reactivation, a treatment known to disrupt memory consolidation is applied. Later (usually at least 24 h after the reactivation session) the memory retention is tested by presenting the CS again (Nader, 2003; Tronson and Taylor, 2007).

On the basis of literature results shown in Tables 1–3, it is clear
 that the amygdala and the hippocampus are the neural structures
 most investigated in this mnemonic phase. This is probably due to

#### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

### Table 1

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Reconsolidation: effects of intra-amygdala post recall treatments on fear memory. The table lists the studies using local administration of pharmacological or genetic treatments to determine the role of amygdala and amygdaloid signaling molecules in fear memories reconsolidation.

Cerebral site	Behavioral paradigm	Treatment	Effect	Reference
Amygdala	Inhibitory avoidance	TTX	Impairment	Prado-Alacalà et al. (2006)
BLA	Inhibitory avoidance	Lidocaine	Impairment	Tzeng et al. (2012)
BLA	Auditory fear conditioning	TTX	Impairment	Sacchetti et al. (2007)
BLA	Auditory fear conditioning	TTX	No effect	Baldi et al. (2008)
BLA	Contextual fear conditioning	TTX	Impairment	Baldi et al. (2008) and Bucherelli et al. (2006)
BLA	Auditory fear conditioning	NMDA antagonist	Impairment	Mamou et al. (2006)
BLA	Auditory fear conditioning	NMDA agonist	Improvement	Lee et al. (2006)
BLA	Auditory fear conditioning	NR2A antagonist	Impairment	Milton et al. (2013)
		NR2B antagonist	No effect	~
		AMPA antagonist	No effect	
LA	Auditory fear conditioning	CP-AMPA receptor blockade	Impairment	Hong et al. (2013)
BLA	Auditory fear conditioning	GR antagonist	Impairment	Jin et al. (2007)
BLA	Inhibitory avoidance	GR antagonist	Impairment	Tronel and Alberini (2007)
LA	Auditory fear conditioning	β-AR antagonist	Impairment	Debiec et al. (2011) and Debiec and LeDoux
				(2004)
		β-AR agonist	Improvement	
LA/BLA	Fear-potentiated startle	CB1 agonist	Impairment	Lin et al. (2006)
,	ī	CB1 antagonist	No effect	
BLA	Contextual fear conditioning	CB1 antagonist	Impairment	Bucherelli et al. <mark>(2006)</mark>
	5	H3 antagonist	No effect	Λ
		Muscarinic antagonist	No effect	
BLA	Inhibitory avoidance	Noradrenaline	No effect	Cammarota et al. (2004)
BLA	Auditory fear conditioning	PKA activator	Improvement	Tronson et al. (2006)
	mantory real conditioning	PKA inhibitor	Impairment	
LA	Auditory fear conditioning	MAPK inhibitor	Impairment	Diaz-Mataix et al. (2011) and Duvarci et al. (2005)
BLA	Contextual fear conditioning	PI-3 <mark>K</mark> inhibitor	Impairment	Kritman and Maroun (2013)
BLA	Inhibitory avoidance	Protein synthesis inhibitor	No effect	Cammarota et al. (2004)
BLA	Auditory fear conditioning	Protein synthesis inhibitor	Impairment	Debiec et al. (2006), Duvarci et al. (2006),
	haditory real conditioning	Totem synthesis minister	impuirment	Duvarci and Nader (2004), Jarome et al. (2012),
				Mamou et al. (2006), Nader et al. (2000),
				Sacchetti et al. (2007) and Wang et al. (2009)
LA	Auditory fear conditioning	Protein synthesis inhibitors	Impairment	Debiec et al. (2010) and Duvarci et al. (2005)
Amygdala	Contextual fear conditioning	Protein synthesis inhibitor	Impairment	Mamiya et al. (2009) and Parsons et al. (2006a)
Amygdala	Auditory fear conditioning	mRNA synthesis inhibitors	No effect	Parsons et al. (2006a)
Amygdala	Contextual fear conditioning	mRNA synthesis inhibitors	No effect	Parsons et al. (2006a)
BLA	Auditory fear conditioning	mRNA synthesis inhibitors	Impairment	Duvarci et al. (2008)
BLA	Inhibitory avoidance	C/EBPβ antisense ODN	Impairment	
LA	Auditory fear conditioning	EGR-1 (ZIF268) antisense ODN	Impairment	Tronel et al. (2005) Maddox et al. (2011)
BLA		CREB inhibition	•	Tronson et al. (2012)
LA	Auditory fear conditioning	elF4F inhibitor	Impairment No effect	Hoeffer et al. (2012)
	Auditory fear conditioning	FGF2		
LA	Auditory fear conditioning		Impairment	Graham and Richardson (2011) Madday and Schofa (2011) and Madday at al
LA	Auditory fear conditioning	HDAC inhibitor	Improvement	Maddox and Schafe (2011) and Maddox et al. (2013, 2014)
		DNMT inhibitor	Impairment	(==::;=:::)
		HAT inhibitor	Impairment	
Amygdala	Auditory fear conditioning	mTOR inhibitor	Impairment	Parsons et al. (2006b)
BLA	Inhibitory avoidance	mTOR inhibitor	Impairment	Jobim et al. (2012)
BLA	Auditory fear conditioning	Actin inhibitor	Impairment	Rehberg et al. (2010)
BLA	Contextual fear conditioning	Actin inhibitor	Impairment	Motanis and Maroun (2012) and Rehberg et al.
ULA.	CONCERCIAL ICAL CONCILIONING		mpanment	motants and maroun [2012] and Renderg et al.

Antisense ODN, antisense oligodeoxynucleotide; AMPA α-amino-3-hydroxy-5-methyl-4-isoxazole propionic acid receptor; β-AR β-adrenergic receptor; BLA, basolateral amygdala; CB1, cannabinoid receptor type 1; C/E=Pβ, CCAAT enhancer-binding protein-β; CP-AMPA, calcium permeable AMPA receptors; CREB, cyclic AMP response element-binding protein; DNMT, DNA methyltransferase; EGR\_1, early growth response gene-1; elF4F, eukaryotic initiation factor 4F; FGF2, fibroblast growth factor 2; GR glucocrticoid receptor; H3, histaminergic receptor type 3; HAT, histone acetyltransferase; HDAC, histone deacetylase; MAPK, mitogen-activated protein kinase; mTOR mammalian target of rapamycin kinase; NMDA, N-methyl-D-aspartate receptor; NR2A, NR2B, NMDA receptors subtype 2A, 2B; Pl-3K, phosphatidylinositol 3 kinase; PKA, protein kinase A; TTX, tetrodotoxin; ZIF268, zinc finger 268.

fear memory reconsolidation. Moreover, the blockade of CP-AMPA receptor immediately after retrieval impairs the reconsolidation process of auditory fear memory.

Noradrenergic transmission is also involved in reconsolidation process; in fact, post-reactivation intra  $\overline{\ }$  BLA administration of  $\beta$ -adrenergic receptor ( $\beta$ -AR) antagonist or agonist reduces or enhances, respectively, fear memory (Debiec et al., 2011; Debiec and LeDoux, 2004). Also post-reactivation blockade of glucocorticoid receptors (GRs) in this neural site disrupts long-term fear memories retention (Jin et al., 2007; Tronel and Alberini, 2007). In our laboratory we studied the BLA cholinergic, histaminergic and cannabinoid systems involvement in contextual fear memory reconsolidation (Bucherelli et al., 2006). The results showed that the cannabinoid system participates in memory maintenance 291 after reactivation, whereas cholinergic and histaminergic neurons 292 do not. Amygdalar cannabinoid receptor type 1 (CB1) involvement 293 was also demonstrated in the fear-potentiated startle reconsolida-294 tion (Lin et al., 2006), although in this case mnemonic impairment 295 followed the activation of these receptors by CB1 agonists and was reverted by a selective CB1 antagonist. The difference between these results and ours is not clear but it could be due to the different fear responses studied (fear-potentiated startle vs. freezing response). Reconsolidation of different fear responses might 300 require either activation or blockade of intra-amygdala CB1 recep-301 tors. Together these results suggest that several neurotransmitter 302 systems within the BLA are critically involved in fear memory 303

### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

### Table 2

Reconsolidation: effects of intra-hippocampus post recall treatments on fear memory. The table lists the studies using local administration of pharmacological or genetic treatments to determine the role of hippocampus and hippocampal signaling molecules in fear memories reconsolidation.

Cerebral site	Behavioral paradigm	Treatment	Effect	Reference
DHC	Inhibitory avoidance	Muscimol	Impairment	Amaral et al. (2007)
DHC	Inhibitory avoidance	TTX	Impairment	Prado-Alacalà et al. (2006)
DHC	Contextual fear conditioning	NMDA antagonist	Impairment	Lee and Hynds (2013)
<b>H</b> ippocampus	Contextual fear conditioning	L-VGCC inhibitor	No effect	Suzuki et al. (2008)
DHC	Inhibitory avoidance	α7nAChR agonist (low footshock intensity)	Improvement	Boccia et al. (2010)
		α7nAChR antagonist (low footshock intensity)	Impairment	X
		$\alpha$ 7nAChR agonist (high footshock intensity)	Impairment	
		$\alpha$ 7nAChR agonist (high footshock intensity)	Impairment	
DHC	Inhibitory avoidance	α7nAChR agonist	Improvement	Blake et al. (2012, 2013)
<b>Hippocampus</b>	Contextual fear conditioning	CB1 antagonist	Improvement	Blake et al. (2012, 2013) De Oliveira Alvares et al. (2008)
	-	CB1 agonist	Impairment	
Hippocampus	Contextual fear conditioning	CB1 inhibitor	No effect	Suzuki et al. <mark>(2008)</mark>
DHC	Inhibitory avoidance	MAPK inhibitor	No effect	Roesler and Quevedo (2009)
DHC	Contextual fear conditioning	MAPK inhibitor	No effect	Lee and Hynds (2013)
DHC	Contextual fear conditioning	IKK inhibitor	Impairment	Lee and Hynds (2013)
DHC	Inhibitory avoidance	IKK inhibitor	Impairment	Boccia et al. (2007)
DHC	Contextual fear conditioning	PI-3 <mark>K i</mark> nhibitor	No effect	Chen et al. $(2005)$
DHC	Inhibitory avoidance	Protein synthesis inhibitor	No effect	Chen et al. (2005) Cammarota et al. (2004) and Taubenfeld et al.
DHC	Inhibitory avoidance	Protein synthesis inhibitor	Impairment	(2001) Power et al. (2006)
DHC	Contextual fear conditioning	Protein synthesis inhibitor	Impairment	Chen et al. $(2005)$ and Debiec et al. $(2002)$ .
				Chen et al. (2005) and Debiec et al. (2002), Frankland et al. (2006), Lee (2008), Lee et al.
				(2004, 2008), Mamiya et al. (2009), Stafford and Lattal (2009) and Suzuki et al. (2008)
DHC	Contextual fear conditioning	mRNA synthesis inhibitors	Impairment	De Oliveira Alvares et al. (2008) and Lee et al.
Dire	contextual real conditioning	ment synthesis minorers	impairment	(2004)
Hippocampus	Inhibitory avoidance	C/EBPβ antisense ODN	No effect	Taubenfeld et al. (2001)
DHC	Inhibitory avoidance	IGF-II	Improvement	Chen et al. (2011)
DHC	Contextual fear conditioning	BDNF antisense ODN	No effect	Chen et al. (2011) Barnes et al. (2012), Lee (2008) and Lee et al. (2004)
DHC	Contextual fear conditioning	ZIF268 antisense ODN	Impairment	Barnes et al. (2012), Kirtley and Thomas
				(2010), Lee (2008) and Lee et al. (2004)
DHC	Contextual fear conditioning	Proteasome inhibitor	No effect	Lee (2008) and Lee et al. (2008)
		Proteasome inhibitor + anisomycin	Anisomycin	N N
		A	effect blockade	
DHC	Contextual fear conditioning	IL-1R antagonist	Impairment	Barnes et al. (2012) and Machado et al. (2010)
Hippocampus	Contextual fear conditioning	NF-kB inhibitor	Impairment	Barnes et al. (2012) and Machado et al. (2010) De la Fuente et al. (2011)
	-	NFAT inhibitor	No effect	Δ
		Calcineurin inhibitor	No effect	
DHC	Inhibitory avoidance	NF-kB inhibitor	Impairment	Boccia et al. (2007)
DHC	Contextual fear conditioning	mTOR inhibitor	Impairment	Gafford et al. (2011)
DHC	Inhibitory avoidance	mTOR inhibitor	Impairment	Jobim et al. (2012)
Hippocampus	Contextual fear conditioning	Actin inhibitor	Impairment	Motanis and Maroun (2012)

 $\alpha$ 7nAChR,  $\alpha$ 7 nicotinic acetylcholine receptor; antisense ODN, antisense oligodeoxynucleotide; BDNF, brain-derived neurotrophic factor; CB1, cannabinoid receptor type 1; C/E=Pβ, CCAAT enhancer-binding protein-β; DHC, dorsal hippocampus; IGF-II, insulin-like growth factor II; IKK, IκB protein kinase; IL-1R, interleukin 1 receptor; L-VGCC, L-type voltage-gated calcium channel; MAPK, mitogen-activated protein kinase; mTOR, mammalian target of rapamycin kinase; NFAT, nuclear factor of activated T-cells; NF-κB, nuclear factor κB; NMDA, N-methyl-p-aspartate receptor; PI-3K, phosphatidylinositol 3 kinase; TTX, tetrodotoxin; ZIF268, zinc finger 268.

### Table 3

Reconsolidation: effects of intra several cerebral sites post recall treatments on fear memory. The table lists the studies using local administration of pharmacological or genetic treatments to determine the role of specific cerebral sites and signaling molecules in fear memories reconsolidation.

Cerebral site	Behavioral paradigm	Treatment	Effect	Reference
mPFC	Contextual fear conditioning	Protein synthesis inhibitor	No effect	Mamiya et al. (2009)
IL-mPFC	Contextual fear conditioning	Muscimol	No effect	Stern et al. (2014)
IL-mPFC	Contextual fear conditioning	PI-3 <mark>K</mark> inhibitor	No effect	Kritman and Maroun (2013)
PL-mPFC	Contextual fear conditioning	Muscimol	Impairment	Stern et al. (2014)
PL-mPFC	Olfactory fear conditioning	α-AR antagonist	Impairment	Do Monte et al. (2013a,b)
Anterior cingulate cortex	Contextual fear conditioning	Protein synthesis inhibitor	No effect	Frankland et al. (2006)
Anterior cingulate cortex	Contextual fear conditioning	Protein synthesis inhibitor	Impairment	Einarsson and Nader (2012)
Anterior cingulate cortex	Olfactory fear conditioning	$\alpha_1$ -AR antagonist	No effect	Do Monte et al. (2013a,b)
ENT	Contextual fear conditioning	TTX	Impairment	Baldi and Bucherelli (2014)
ENT	Inhibitory avoidance	Protein synthesis inhibitor	No effect	Cammarota et al. (2004)
		Noradrenaline	No effect	X · · ·
Perirhinal cortex	Auditory fear conditioning	TTX	No effect	Sacchetti et al. (2007)
		Protein synthesis inhibitor	No effect	Λ .
Cerebellar vermis	Auditory fear conditioning	TTX	Impairment	Sacchetti et al. (2007)
		Protein synthesis inhibitor	Impairment	Λ
Nucleus basalis magnocellularis	Auditory fear conditioning	TTX	No effect	Baldi et al. <b>(2008)</b>
Nucleus basalis magnocellularis	Contextual fear conditioning	TTX	No effect	Baldi et al. 2008)

 $\alpha$ -AR,  $\alpha$ -adrenergic receptor;  $\alpha_1$ -AR,  $\alpha_1$ -adrenergic receptor; ENT, entorhinal cortex; IL-mPFC, infralimbic subregion of the medial prefrontal cortex; mPFC, medial prefrontal cortex; PI-3K, phosphatidylinositol 3 kinase; PL-mPFC, prelimbic subregion of the medial prefrontal cortex; TTX, tetrodotoxin.

reconsolidation, and also show that this process cannot be considered a recapitulation of consolidation, although some mechanisms in BLA are in common.

### 2.1.2. Protein kinases

The activation/inhibition of several neurotransmitter systems in the amygdala during fear memory reconsolidation is thought to lead, either directly or indirectly, to the activation of downstream signaling cascades. Two protein kinases are of particular interest: protein kinase A (PKA) and mitogen-activated protein kinase (MAPK). These kinases were shown to contribute to fear memory consolidation engaging cellular processes necessary for long-term synaptic plasticity and memory formation. PKA and MAPK are required for conditioned auditory fear reconsolidation into the BLA. In this brain structure, PKA activation enhances reconsolidation processes (Tronson et al., 2006), whereas PKA or MAPK inhibition impairs memory reconsolidation (Diaz-Mataix et al., 2011; Doyere et al., 2007; Duvarci et al., 2005; Tronson et al., 2006). Moreover, auditory fear memory reconsolidation impairment induced by MAPK inhibition was obtained using both discrete CS and US for reactivating the mnemonic trace (Diaz-Mataix et al., 2011; Doyere et al., 2007). The consequent successive loss of reinstatement suggests that this loss of fear memory, and its neurophysiological correlate in the BLA induced by the MAPK inhibitor after (US or CS) reactivation, does not reflect a retrieval blockade. Also, the phosphatidylinositol-3 kinase (PI-3K) and its target, AKT/PKB (protein serine/threonine kinase), are critical for memory reconsolidation. In fact, Kritman and Maroun (2013) reported that PI-3K inhibition into the BLA before retrieval of a contextual fear task impairs reconsolidation of this memory. Because PI-3K and AKT/PKB are upstream targets of the mammalian target of rapamycin (mTOR) pathway, these results prove that PI-3K-AKT/PKB-mTOR pathway has a crucial role in fear memory reconsolidation at least into the BLA.

### 2.1.3. Gene expression and protein synthesis

PKA and MAPK act directly or indirectly activating several transcription factors, such as CREB (cyclic AMP-response element binding protein) and zif268 (zinc finger 268) that initiate gene transcription. Some of these transcription factors within the BLA are implicated in fear memory reconsolidation. For example, CREB inhibition or ICER (inducible CREB early repressor) overexpression into the BLA induced impairment of auditory fear memory reconsolidation (Tronson et al., 2012). Moreover, the inhibition of CREB activity did not disrupt memory retrieval. Thus, these results support the idea that disruption of reconsolidation is due to post-retrieval storage failure and not to retrieval impairment (Alberini, 2008; Hardt et al., 2009; Nader et al., 2000; Riccio et al., 2002).

Another transcription factor that is believed to be critical for regulating the transcription of late-response genes that promote functional and/or structural changes underlying memory formation is zif268 (also known as EGR-1). Regulation of zif268 mRNA in the BLA following auditory and contextual fear memory retrieval (Hall et al., 2001; Maddox et al., 2011) suggests that zif268 is critical for the reconsolidation process in the BLA.

Targeted disruption of the transcription factor CCAAT enhancer binding protein  $\beta$  (C/EBP $\beta$ ) in the BLA impairs reconsolidation of fear memory, specifically of inhibitory avoidance. Within the BLA C/EBPB appears to be required for reconsolidation but not for consolidation of this mnemonic task (Tronel et al., 2005). This result has been considered as an example of dissociation between the two processes. In other words, the different C/EBPβ requirement in the BLA during these memory phases can be used to dissociate the two processes both at the anatomical and molecular level.

Gene transcription can also be controlled by epigenetic mechanisms and recent studies have focused on those that might be involved in memory reconsolidation. Epigenetic mechanisms include modifications in chromatin structure and DNA methylation. 360 Chromatin consists of DNA packaged around a core of eight histones 370 and it is post-translationally regulated by acetylation of histones on 371 their N-terminal tails via histone acetyltransferases (HATs). This 372 induces the relaxation of chromatin structure, leading to enhanced 373 transcription, and can be reversed by histone deacetylases (HDACs) 374 (Levenson and Sweatt, 2005). On the other hand, DNA methylation 375 is associated with transcriptional repression which is catalyzed by 376 DNA methyltransferases (DNMTs) (Levenson and Sweatt, 2005). 377 Histone acetylation of chromatin is thought to positively regu-378 late transcription, whereas DNA methylation has a negative effect 379 on transcription regulation. Recently, these two processes were 380 shown to be crucial for fear memories reconsolidation in the amyg-381 dala. Maddox and coworkers (Maddox and Schafe, 2011; Maddox 382 et al., 2014) reported that intra-LA infusion of inhibitors of HDAC 383 and DNMT activity enhanced and impaired auditory fear memory 384 reconsolidation, respectively. Moreover, the same authors showed 385 that p300/CBP histone acetyltransferase activity within the BLA is 386 critical for reconsolidation of auditory fear conditioning (Maddox 387 et al., 2013), as intra-LA infusion of an inhibitor of the p300/CBP 388 HAT impaired memory reconsolidation.

Post-retrieval inhibition of protein synthesis has been one of 390 the most frequently used treatments to investigate the nature of 391 memory reconsolidation. This use of protein synthesis inhibitors 392 relies on the fact that protein synthesis is considered a marker 393 of consolidation processes, necessary to render structural cellular 30/ changes permanent, and of the involvement of a neural region in 305 mnemonic phase. Most of these studies showed that the local injec-306 tions of protein synthesis inhibitors (anisomycin or cycloheximide) 397 into the BLA after retrieval of a consolidated auditory or contex-308 tual fear memory impaired the original memory (Debiec et al., 399 2006, 2010; Duvarci et al., 2006; Duvarci and Nader, 2004; Duvarci 400 et al., 2005; Jarome et al., 2012; Mamiya et al., 2009; Mamou 401 et al., 2006; Nader et al., 2000; Parsons et al., 2006a; Sacchetti 402 et al., 2007; Wang et al., 2009). Thus, these results provided evi-403 dence that fear memories, once reactivated, must undergo protein 404 synthesis-dependent reconsolidation in the BLA to be maintained 405 for subsequent retrieval. Moreover, this reconsolidation process 406 has a temporal window during which blockade of protein synthesis 407 is effective. 408

There are also negative results. Cammarota et al. (2004) reported that the intra-BLA infusion of anisomycin performed before or after 410 a reactivation session of an inhibitory avoidance task does not affect 411 subsequent memory retention; accordingly it does not seem there 412 is a retrieval-induced, protein synthesis-dependent process that 413 would cause reconsolidation of this fear memory. 414

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During reconsolidation does protein synthesis depend on 415 already existing mRNAs or on synthesis of new mRNAs in the 416 BLA? On this point, contradictory results were obtained; in fact, 417 Parsons et al. (2006a) have shown that auditory and contextual 418 fear memory reconsolidation is independent on mRNA synthesis 419 in the amygdala, whereas according to Duvarci et al. (2008) this 420 process requires de novo mRNA synthesis in this neural structure. 421 Likely, as underlined by the authors, the different results may be 422 ascribed to procedural differences because it is possible that even 423 small changes in experimental procedures can alter the molecu-424 lar mechanisms engaged (Tronson and Taylor, 2007). There are at 425 least two forms of protein synthesis: the primary mode of trans-426 lation initiation requires formation of a multi-protein complex of 427 eukaryotic initial factors (eIFs) bound to the 5' methylated-GTP cap 428 of target mRNAs. Specifically, the interaction between eIF4E and 429 eIF4G facilitates eIF4A RNA helicase activity, recruitment of the 40 430 S ribosomal subunit, scanning, and peptide elongation. Molecules 431 that block the formation of eIF4F (eIF4E<sub>4</sub>+eIF4G+eIF4A), such as 432 the endogenous regulator 4E-binding protein, which binds to 433

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### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

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and sequesters eIF4E, therefore effectively inhibits cap-dependent 434 translation. Likewise, the small molecule, 4EGI-1, which selectively 435 disrupts eIF4E-eIF4G interactions (eIF4F formation) in vitro, also 436 inhibits cap-dependent translation. The second route by which 437 mRNAs can be translated occurs via internal ribosomal entry sites 438 (IRES), which are unaffected by disruptions to the 5' cap trans-430 lation machinery, such as blockade of eIF4E-eIF4G interactions. 440 Very little is known about the specific mechanistic constraints on 441 the phases of the memory processes. By microinfusing 4EGI-1 into 442 the LA, the authors investigated the role of cap-dependent trans-443 lation and eIF4F formation in reconsolidation of the cued (tone) 444 fear conditioning. 4EGI-1 impaired consolidation but not recon-445 solidation. Thus, these two memory processes require different 446 translational control mechanisms. In other words, consolidation 447 is dependent on eIF4E-eIF4G interactions or required for cap-448 dependent protein synthesis; instead, reconsolidation does not 449 seem to require cap-dependent protein synthesis, although it is 450 possible that eIF4E\_eIF4G interactions are increased during this 451 memory phase in a temporal window outside those considered in 452 the experiments (Hoeffer et al., 2011). 453

Neurons protein synthesis is regulated at the translational level through phosphorylation of several intracellular targets. In particular, the signaling pathway controlled by mTOR kinase regulates protein translation by controlling the phosphorylation state of the eIF4E-binding protein 1 (4E-BP1) and p70s6 kinase (p70s6K) (Raught et al., 2001). Post-retrieval intra-BLA infusion of rapamycin, an inhibitor of the mTOR pathway, disrupts reconsolidation of auditory fear memory after retrieval (Parsons et al., 2006b). As considerable evidence shows that many of the effects of mTOR on plasticity are localized to dendrites, this result seems to suggest that mTOR pathway may be involved in regulating the local protein synthesis that supports memory reconsolidation (Parsons et al., 2006b). This same signaling pathway is thought to be also involved in inhibitory avoidance as rapamycin impaired long-term retention of this memory when given before or immediately after retrieval into the BLA (Jobim et al., 2012).

During memory formation, structural changes at synapses occur 470 and transcriptional and translation processes might serve to re-471 stabilize these changes and to maintain the memory trace. These 472 synaptic alterations may involve re-arrangement of the actin 473 474 cytoskeleton. Actin filaments are critically involved in several synaptic functions, such as control of neurotransmitter exocytosis 475 (Morales et al., 2000), vesicles recycling (Shupliakov et al., 2002), 476 trafficking of neurotransmitters receptors and structural modifica-477 tion of post-synaptic spines (Honkura et al., 2008; Zhou et al., 2001). 478 Intra-BLA injection of toxin cytochalasin D, which depolymerizes 479 actin filaments, blocks contextual fear memory reconsolidation 480 (Motanis and Maroun, 2012). Similar results were obtained using 481 the death cap toxin phalloidin that arrests actin filaments (Rehberg 482 et al., 2010). Intra-BLA application of phalloidin impairs reconsol-483 idation of auditory and contextual fear memory when performed 484 30 min after reactivation session; the same treatment performed 485 6h after reactivation impairs reconsolidation of auditory mem-486 ory trace, but not reconsolidation of contextual ones (Rehberg 487 et al., 2010). Thus, these results suggest a crucial role of actin re-488 arrangement in reconsolidation process of fear memories. 489

### 2.2. Hippocampus

Whereas the amygdala is crucial for fear memory associated to either a discrete CS and contextual CS, the hippocampus is necessary for contextual fear memory (inhibitory avoidance included), but not for auditory fear conditioning. Several studies used inactivating agents (such as TTX and muscimol, GABA<sub>A</sub> receptor agonist) which depress neuronal excitability to study the hippocampal role in fear memory reconsolidation (Table 2). The results showed that post-reactivation infusion of these pharmacological agents into the dorsal hippocampus disrupted retention of inhibitory avoidance memory (Amaral et al., 2007; Prado-Alacalà et al., 2006). However, the deficit was temporary as it reversed spontaneously with time in the absence of multiple testing (Amaral et al., 2007) and it was attenuated progressively with repeated retention testing (Prado-Alacalà et al., 2006).

### 2.2.1. Neurotransmitter systems

Hippocampal-dependent fear memories reconsolidation requires several neurotransmitter systems. The contextual fear reconsolidation is impaired by the injection of NMDA antagonist in the hippocampus 15 min before the reactivation session (Lee and Hynds, 2013) demonstrating the importance of this receptor in this mnemonic phase. A critical role in reconsolidation of inhibitory avoidance is played by the hippocampal  $\alpha$ 7 nicotinic acetylcholine receptor ( $\alpha$ 7nAChR). Specifically, hippocampal  $\alpha$ 7nAChR activation by the agonist choline after reactivation of an inhibitory avoidance memory, impaired subsequent retention test in mice trained with a high footshock intensity, whereas the memory retention was improved in mice trained with a low footshock intensity (Boccia et al., 2010). However, Blake et al. (2012, 2013) observed memory reconsolidation improvement 519 following α7nAChR agonist administration also using high shock intensity. On the contrary, intra-hippocampus injection of an 521 α7nAChR antagonist impaired memory reconsolidation regardless 522 of footshock intensity (Boccia et al., 2010).

Contradictory results were obtained about CB1 receptor role 524 in contextual fear memory reconsolidation. Suzuki et al. (2008) 525 found that hippocampal CB1 antagonist administration immedi-526 ately after a brief re-exposure to training context had no effect 527 on memory retention. However, when the CB1 antagonist was 528 co-administered with anisomycin after context re-exposure, it 529 protected contextual memory against the amnesic effects of ani-530 somycin. The same authors showed similar results using a L-type 531 voltage-gated calcium channel (L-VGCC) antagonist: the blockade 532 of this ionic channel had no effect per se on contextual memory 533 reconsolidation, but its co-administration with protein synthe-534 sis inhibitor anisomycin prevented the disruption of reactivated 535 memory. Thus, in the hippocampus, CB1 and L-VGCC mediate 536 destabilization of contextual fear memory which occurs following 537 the reactivation session. On the other hand, De Oliveira Alvares et al. 538 (2008) reported that a CB1 antagonist infused intra-hippocampus 539 after a reactivation session caused facilitation of contextual mem-540 ory reconsolidation. The local administration of a CB1 agonist 541 caused disruption of this mnemonic process and this effect was 542 abolished by the combined administration of a CB1 agonist and 543 antagonist. 544

### 2.2.2. Protein kinases

Unlike what has been observed in the in amygdala, PI-3K inhibitors injection into the hippocampus has no effect on contextual memory reconsolidation (Chen et al., 2005).

Although fear memory consolidation and reconsolidation show an overlap concerning some molecular mechanisms, independent cellular processes were reported within the hippocampus in the two phases of contextual fear memorization. Dissociation was observed for the requirement of MAPK and IkB kinase (IKK) (Lee and Hynds, 2013). Administration of MAPK inhibitor into the dorsal hippocampus did not affect either contextual nor inhibitory avoidance reconsolidation, but impaired their initial consolidation (Lee and Hynds, 2013; Roesler and Quevedo, 2009). Instead, inhibition of hippocampal IKK induced impairment of memory reconsolidation without affecting consolidation (Boccia et al., 2007; Lee and Hynds, 2013). 560

#### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

### 2.2.3. Gene expression and protein synthesis

A double dissociation was shown between the transcription factors BDNF (brain-derived neurotrophic factor) and zif268; whereas contextual fear memory consolidation depends on BDNF (Barnes et al., 2012; Lee, 2008; Lee et al., 2004), its reconsolidation requires zif268 (Barnes et al., 2012; Kirtley and Thomas, 2010; Lee, 2008; Lee et al., 2004). Finally, dissociation was observed in the hippocampal C/EBPβ. Using an inhibitory avoidance task, Taubenfeld et al. (2001) reported that this transcription factor in the hippocampus is required for consolidation of a new inhibitory avoidance memory, but nor for a reactivated fear memory.

Using microarray analysis, Barnes et al. (2012) showed that in the hippocampus activation of some genes is shared between consolidation and reconsolidation of contextual fear memory. These genes, however, are regulated in opposite directions. In particular, among the shared genes, there are those associated with pro-inflammatory cytokine pathway that appear to be downregulated during consolidation and upregulated during reconsolidation of contextual fear memory. Also, the injection of an interleukin 1a (IL-1a) receptor antagonist into the hippocampus immediately after retrieval reduced retention of the recalled contextual memory indicating that in the hippocampus the contextual fear memory reconsolidation depends on IL-1a receptor pathway. However, there is no direct experimental evidence about IL-1a antagonism on contextual fear conditioning consolidation. Involvement of the cytokines in contextual fear memory reconsolidation was shown by Machado et al. (2010). These authors reported that intrahippocampus administration of interleukin 1 $\beta$  up to 30 min after reactivation session decreased subsequent memory retention. This effect was reversed by  $\alpha$ -melanocyte-stimulating hormone that had no effect per se on contextual fear memory reconsolidation.

A putative C/EBPβ gene is the insulin-like growth factor II (IGF-II) which has relatively high concentration within the hippocampus. Chen et al. (2011) investigated the functional role of this growth factor in memory reconsolidation. They showed that hippocampal injection of IGF-II after retrieval of inhibitory avoidance memory enhanced subsequent memory retention. However, whether the treatment was performed immediately post-retrieval, two weeks after training, did not induce memory enhancement during retention testing. Thus, memory improvement induced by hippocampal IGF-II occurs only when the temporal window during which inhibitory avoidance memory undergoes reconsolidation.

Cellular imaging has shown that some immediate early genes are activated after retrieval of a previously consolidated memory. In the hippocampus, the retrieval of contextual fear memory is followed by c-Fos and JunB activation, while c-Jun or JunD are not activated (Strekalova et al., 2003). Other IEGs considered were serum- and glucocorticoid-induced kinase 3 (SGK3) and nerve growth factor inducible gene B (NGFI-B). Among these IEGs, SGK3 is upregulated both after training and retrieval of contextual fear in the hippocampus, whereas NGFI-B is regulated only during consolidation (Von Hertzen and Giese, 2005).

Post-retrieval inhibition of protein synthesis has been one of the most used treatments to analyze hippocampus-dependent fear memories reconsolidation, such as just contextual fear conditioning and inhibitory avoidance. It was demonstrated that intra-hippocampus anisomycin injection performed after reactivation of contextual fear memory caused a mnemonic impairment at subsequent retention test (Chen et al., 2005; Debiec et al., 2002; Frankland et al., 2006; Lee, 2008; Lee et al., 2004, 2008; Mamiya et al., 2009; Stafford and Lattal, 2009; Suzuki et al., 2008). Thus, it was concluded that the contextual fear memory stored in the hippocampus undergoes a protein synthesis-dependent reconsolidation process whenever it is reactivated. In other words, hippocampal memory reconsolidation depends on de novo protein synthesis. In contrast, it was shown that injections of anisomycin into the hippocampus were ineffective in blocking reconsolidation 627 of inhibitory avoidance (Cammarota et al., 2004; Taubenfeld et al., 628 2001) or the blockade is temporary (Power et al., 2006). These 620 results provide evidence that hippocampal protein synthesis is 630 not requested for inhibitory avoidance reconsolidation. Moreover, 631 they also raised the hypothesis that reconsolidation, as a protein 632 synthesis-dependent process, does not occur in this neural site. 633 However, it must be underlined that inhibitory avoidance memory 63/ is impaired by systemic administration of anisomycin performed 635 following memory recall (Taubenfeld et al., 2001). The different hip-636 pocampal involvement in the reconsolidation of the two paradigms 637 might be due to the different requirements of the tasks. In fact, 638 the inhibitory avoidance is much more complex than classical con-639 textual fear conditioning and requires an instrumental response. 640 The transience of retrieval impairment has been used by some 641 authors to argue against reconsolidation process on the basis that 642 its blockade does not produce the same effects as blocking consol-643 idation (Power et al., 2006). However, the reversibility of amnesia 644 does not necessarily constitute evidence against the reconsolida-645 tion hypothesis because these same studies have shown that this 646 effect is dependent on memory reactivation (Amaral et al., 2007; 647 Power et al., 2006).

De novo protein synthesis which occurs during hippocampal contextual fear memory reconsolidation appears to depend on de novo mRNA synthesis. Local injections of mRNA synthesis inhibitors after re-exposure trial impair retention of this memory (De Oliveira Alvares et al., 2008; Lee et al., 2004).

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The memory reconsolidation consists of two phases: a desta-654 bilization process and re-stabilization ones (Hong et al., 2013; 655 Lee, 2008). It has been hypothesized that during memory re-656 stabilization/reconsolidation process, removal of existing proteins 657 and incorporation of new proteins occur. For this purpose some 658 authors investigated the involvement of hippocampal proteasome 659 system (the main cellular mechanism controlling protein turnover) 660 in the contextual fear memory destabilization/re-stabilization after 661 retrieval. The results showed that hippocampal injection of the 662 proteasome inhibitor βlac immediately after retrieval/reactivation 663 session has no effect on subsequent contextual memory retention. 664 However, the co-administration of Blac and anisomycin prevented 665 memory reconsolidation impairment induced by anisomycin alone 666 (Lee, 2008; Lee et al., 2008). Thus, proteasome-dependent pro-667 tein degradation after memory retrieval destabilizes preexisting contextual fear memory which then undergoes reconsolidation process (Lee et al., 2008). Moreover, inhibition of memory desta-670 bilization may maintain the strength of a previously acquired 671 memory supporting the concept that memory reconsolidation 672 allows the strengthening of memory (Lee, 2008). 673

As for the amygdala, it was demonstrated that hippocampal 674 protein synthesis is regulated, at least in part, by mTOR pathway. 675 Intra-hippocampus administration of either transcription factor 676 NF- $\kappa$ B(nuclear factor- $\kappa$ B) inhibitor (Boccia et al., 2007; De la Fuente 677 et al., 2011) or transcriptional inhibitor rapamycin (Gafford et al., 678 2011; Jobim et al., 2012) impaired reconsolidation of contextual 679 and inhibitory avoidance memories (Boccia et al., 2007; De la 680 Fuente et al., 2011; Gafford et al., 2011; Jobim et al., 2012). Probably, 681 the hippocampal mTOR pathway is not activated by PI-3K since it 682 was reported that PI-3K inhibitors injection into the hippocampus 683 has no effect on contextual memory reconsolidation (Chen et al., 684 2005). Instead, it was shown that within this neural site translation 685 control through mTOR pathway is also crucial for consolidation of 686 contextual and inhibitory avoidance memories (Boccia et al., 2007; 687 Gafford et al., 2011). Therefore, in the hippocampus there is an over-688 lap between molecular mechanisms underlying the fear memory consolidation and reconsolidation.

Structural changes of synapses also occur during hippocampusdependent memories reconsolidation. The hippocampal actin

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### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

rearrangement plays an important role since the infusion of the actin rearrangement antagonist after reactivation session blocks subsequent memory retention (Motanis and Maroun, 2012).

### 696 2.3. Cortex and other neural sites

Recently, researchers have investigated the potential role of 607 other brain regions in fear memories reconsolidation (Table 3). 698 Among these, several cortical regions have been considered. The 699 medial Prefrontal Cortex (mPFC) appears to be involved in cued fear 700 memory reconsolidation, whereas there are contradictory results 701 about its role in contextual fear reconsolidation. Using an olfac-702 tory fear conditioning paradigm Do Monte et al. (2013b) studied 703 the involvement of two mPFC sub-regions, the prelimbic region 704 (PL) and the anterior cingulate cortex (ACC). The results showed 705 that post-retrieval blockade of  $\alpha_1$ -adrenergic receptor within PL-706 mPFC impaired fear memory reconsolidation, whereas the same 707 treatment performed intra-ACC did not affect subsequent mem-708 ory retention (Do Monte et al., 2013b). Concerning contextual fear memory reconsolidation, infusion of muscimol into the PL-710 711 mPFC, but not into IL-mPFC, after reactivation of this memory trace disrupts its subsequent retention (Stern et al., 2014). More-712 over, contextual fear memory reactivation is followed by increased 713 expression of zif268/EGR1 in the PL-mPFC but not in IL-mPFC 714 (Stern et al., 2014). In contrast, Mamiya et al. (2009) reported 715 that (i) CREB-mediated gene expression is not activated in the 716 mPFC (both PL and infralimbic regions) when contextual mem-717 ory is reconsolidated, and (ii) blocking protein synthesis in the 718 mPFC does not affect reconsolidation of this mnemonic trace. Also, 719 the post-reactivation blockade of protein synthesis in the ACC 720 has no effect on contextual memory reconsolidation (Frankland 721 et al., 2006). However, a recent study has shown that injection 722 of anisomycin into the ACC immediately post-reactivation ses-723 sion blocked reconsolidation of this engram (Einarsson and Nader, 724 2012). The differences between these studies may be attributed to 725 the different parameters employed. Finally, Kritman and Maroun 726 (2013) have demonstrated that PI-3K inhibition in the IL-mPFC 727 before retrieval of contextual fear memory does not influence either 728 retrieval or reconsolidation of the mnemonic trace. Thus, together 729 these results suggest that fear reconsolidation occurs in the mPFC, 730 although the specific subregion recruited may depend on the con-731 ditioned stimulus. 732

In our laboratory, we have recently investigated the Entorhinal Cortex (ENT) role during reconsolidation of fear memories. We found that TTX inactivation of the ENT immediately after a brief reactivation session impairs reconsolidation of contextual fear conditioning (Baldi and Bucherelli, 2014). This result does not confirm those by Cammarota et al. (2004) in inhibitory avoidance which show that infusion of anisomycin or noradrenaline (a well-known retrieval enhancer) in this cortical site performed 15 min or 3 h after the reactivation session does not affect subsequent memory retention.

Another cortical site whose potential involvement in reconsolidation was studied is the perirhinal cortex. This neural site is not involved in auditory fear reconsolidation, as local TTX inactivation or anisomycin injection do not alter retrieved fear trace (Sacchetti et al., 2007). The same authors also investigated the role of the cerebellum (more specifically the cerebellar vermis). TTX cerebellar vermis blockade or cerebellar anisomycin injections induced amnesia if performed immediately after the retrieval of auditory fear memory. This effect did not recover over time, even after a reminder footshock administration. Moreover, using a stronger conditioning, the fear memory reconsolidation was affected by the combined but not independent cerebellar and amygdala blockade. Together these results suggest that the cerebellar vermis is a critical neural sites for fear memory reconsolidation and it may support this process even in the absence of the amygdala (Sacchetti et al., 2007).

In our laboratory the nucleus basalis magnocellularis (NBM) role in fear memory reconsolidation was studied. This interest is derived from our previous demonstration that this neural site, which constitutes the main source of cholinergic projections to the cortex and amygdala, is involved in the consolidation of both auditory and contextual engrams in fear conditioning (Baldi et al., 2007). We found that the NBM is not involved in the post-reactivation phase of fear memories. The TTX NBM inactivation performed immediately postreactivation is not followed by an impairment of either acoustic CS or contextual memory trace (Baldi et al., 2008). Thus, unlike the consolidation phase, the relationship between NBM and amygdala might not be equally important during the reconsolidation ones.

### 3. Brain structures involved in fear memory extinction

As mentioned above, fear memory retrieval can initiate another process in addition to the mnemonic trace reconsolidation: memory extinction. Operationally, the engram reactivation is very similar to an extinction session. However, the results are quite different. While the reconsolidation allows, at least partially, the strengthening of the original memory, the extinction weakens its expression. An important factor for the fate of the engram following its retrieval is the duration of re-exposure to the conditioned stimulus in the absence of reinforcement. If the re-exposure is short the reconsolidation process will be triggered, while longer reexposures will induce extinction (Debiec et al., 2002; De la Fuente et al., 2011; Eisenberg et al., 2003; Lee et al., 2006; Pedreira and Maldonado, 2003; Suzuki et al., 2004).

Experimentally, fear memory extinction can be studied by exposing a previously conditioned subject to the repeated nonreinforced presentation of CS. Cued fear extinction is obtained through the repeated exposure to the cue (tone, visual stimulus or odor) in a new environment, whereas contextual fear extinction is obtained by repeatedly presenting the training context. The subsequent extinction memory can be tested either in presence of the discrete CS or in the acquisition context, respectively. Because in the rodent it is thought that extinction is mainly a new learning, two phases have been distinguished: acquisition and consolidation (Myers and Davis, 2007; Pape and Pare, 2010; Quirk and Mueller, 2008). Depending on the phase studied, the treatment will be applied or before extinction training (acquisition), or immediately after extinction training (consolidation). Several evidence indicates that the amygdala, hippocampus and medial prefrontal cortex play a central role in fear extinction (Tables 4–9). Nevertheless, there is no consensus about the specific role of each brain region in the two phases (acquisition, consolidation) of memory extinction.

### 3.1. Amygdala

The BLA seems to be critical in fear extinction (Tables 4 and 7). 803 Using cued and contextual fear conditioning paradigms, it was 804 reported that muscimol-induced inactivation of the BLA performed 805 before extinction training causes impairment of fear memory 806 extinction (Herry et al., 2008; Holmes et al., 2013; Laurent et al., 807 2008; Laurent and Westbrook, 2008, 2010; Sierra-Mercado et al., 808 2011). Thus, the neuronal activity in the BLA seems to be necessary 809 for acquisition of fear extinction. However, contradictory results 810 were obtained when the BLA inactivation was induced immediately 811 after extinction training. Whereas Sierra-Mercado et al. (2011) 812 reported no effect on retention of auditory fear extinction, Akirav 813 et al. (2006) found that intra-BLA muscimol infusion performed 814 immediately after a short extinction training, but not after a long 815 one, facilitates the auditory fear extinction retention. These authors 816

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E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

### Table 4

Extinction acquisition: effects of intra-amygdala pre-extinction training treatments on fear memory extinction. The table shows the studies using local administration of pharmacological or genetic treatments to determine the role of the amygdala and amygdaloid signaling molecules in fear memories extinction acquisition.

Cerebral site	Behavioral paradigm	Treatment	Effect	References
BLA/BA	Auditory fear conditioning	Muscimol	Impairment	Herry et al. (2008) and Sierra-Mercado et al. (2011)
BLA	Cued fear conditioning (CS: light)	Muscimol	Impairment	Holmes et al. (2013) and Laurent and Westbrook (2010)
BLA	Auditory fear conditioning	Muscimol	No effect	Akirav et al. (2006)
BLA	Contextual fear conditioning	Muscimol	Impairment	Laurent et al. (2008) and Laurent and Westbrook (2008)
BLA	Contextual fear conditioning	BZs	Impairment	Hart et al. (2009, 2010)
BLA	Fear-potentiated startle	NMDA agonist	Improvement	Davis (2002), Mao et al. (2006) and Walker et al. (2002)
BLA	Auditory fear conditioning	NMDA agonist	Improvement	Lee et al. (2006)
BLA	Contextual fear conditioning	NMDA agonist	Impairment	Bolkan and Lattal (2014)
BLA	Fear-potentiated startle	NMDA antagonist	Impairment	Davis (2002), Falls et al. (1992), Lin et al. (2003b) and Mao et al. (2006)
BLA	Inhibitory avoidance	NMDA antagonist	Impairment	Myskiw et al. (2010)
BLA	Contextual fear conditioning	NMDA antagonist	Impairment	Laurent et al. (2008) and Lee and Kim (1998)
BLA	Auditory fear conditioning	NMDA antagonist	Impairment	Lee and Kim (1998), Kwapis et al. (2014) and
				Zimmerman and Maren (2010)
BLA	Cued fear conditioning (CS: light)	NMDA antagonist	Impairment	Holmes et al. (2013) and Lee and Kim (1998)
CEA	Auditory fear conditioning	NMDA antagonist	No effect	Zimmerman and Maren (2010)
LA	Auditory fear conditioning	NR2B antagonist	Impairment	Sotres-Bayon et al. (2007, (2009)
BLA	Cued fear conditioning (CS: light)	NR2B antagonist	Impairment	Holmes et al. (2013)
BLA	Auditory fear conditioning	L-VGCC antagonists	Impairment	Davis and Bauer (2012)
BLA	Contextual fear conditioning	NR2B antagonist	Impairment	Laurent et al. (2008) and Laurent and Westbrook (2008)
Amygdala	Contextual fear conditioning	AMPA potentiator	Improvement	Zushida et al. (2007)
BLA	Auditory fear conditioning	AMPA potentiator blockade	Impairment	Kim et al. (2007a,b)
BLA	Fear-potentiated startle	AMPA antagonist	No effect	Falls et al. (1992)
BLA	Auditory fear conditioning	AMPA antagonist	No effect	Zimmerman and Maren (2010)
BLA	Auditory fear conditioning	AMPA antagonist	Impairment	Kwapis et al. (2014)
CEA	Auditory fear conditioning	AMPA antagonist	No effect	Zimmerman and Maren (2010)
LA	Auditory fear conditioning	mGluR1 antagonist	Impairment	Kim et al. (2007a)
BLA	Fear-potentiated startle	CB1 agonist	No effect	Kuhnert et al. (2013)
		CB1 antagonist	No effect	
BLA	Inhibitory avoidance	CB1 agonist	No effect	Ganon-Elazar and Akirav (2009)
		CB1 antagonist	Impairment	
BLA	Auditory fear conditioning	NPS	Improvement	Jungling et al. <mark>(2008)</mark>
DIA	From motortisted startle	NPS inhibitor	Impairment	Version at al. (2000)
BLA	Fear-potentiated startle	GR agonist	Improvement	Yang et al. (2006)
BLA	Fear-potentiated startle	GR antagonist MAPK inhibitor	Impairment Impairment	Davis (2002), Lin et al. (2003b) and Lu et al. (2001)
BLA	Auditory fear conditioning	MAPK inhibitor	Impairment	Herry et al. (2006)
BLA	Inhibitory avoidance	PKA inhibitor	Impairment	Myskiw et al. (2010)
BLA	Inhibitory avoidance	CaMKII inhibitor	Impairment	Myskiw et al. (2010)
BLA	Fear-potentiated startle	PI-3 <mark>K</mark> inhibitor	Impairment	Lin et al. (2003b), Mao et al. (2006) and Yang and Lu (2005)
BLA	Fear-potentiated startle	Calcineurin inhibitor	Impairment	Lin et al. (2003a)
BLA	Fear-potentiated startle	Protein synthesis inhibitor	Impairment	Lin et al. (2003b) and Yang and Lu (2005)
BLA	Fear-potentiated startle	mRNA synthesis inhibitor	No effect	Lin et al. (2003b)
BLA	Fear-potentiated startle	mRNA synthesis inhibitor	Impairment	Yang and Lu (2005)
BLA	Auditory fear conditioning	CREB viral vectors	No effect	Tronson et al. (2012)
BLA	Fear-potentiated startle	BDNF/TrkB viral inhibitor	Impairment	Chhatwal et al. (2006)
		PSA-NCAM cleavage	Improvement	Markram et al. (2007)

AMPA, α-amino-3-hydroxy-5-methyl-4-isoxazole propionic acid receptor; BA, basal amygdala; BDNF, brain-derived neurotrophic factor; BLA, basolateral amygdala; BZs, benzodiazepines; CaMKII<sub>A</sub>Ca<sup>2+</sup>/calmodulin-dependent protein kinase II; CB1, cannabinoid receptor type 1; CEA, central amygdala; CREB<sub>A</sub> cyclic AMP response element-binding protein; GR, glucocorticoid receptor; LA, lateral amygdala; L-VGCC<sub>A</sub>L-type voltage-gated calcium channel; MAPK<sub>A</sub> mitogen-activated protein kinase; mGluR1, metabotropic glutamate receptor subtype 1; NMDA, N-methyl-p-aspartate receptor; NPS<sub>A</sub> neuropeptide S; NR2B, NMDA receptor subtype 2B; PI-3K, phosphatidylinositol 3 kinase; PKA<sub>A</sub> protein kinase A; PSA-NCAM<sub>A</sub> polysialylated neural cell adhesion molecule; TrkB<sub>4</sub> tyrosine kinase B receptor.

concluded that the BLA is involved in extinction consolidation and that GABA<sub>A</sub> transmission facilitates this specific phase of extinction process. Different results were obtained in contextual fear extinction as well. Laurent and Westbrook (2008) showed that intra-BLA post-extinction injection of muscimol impairs extinction retention of this memory; on the contrary, no effect was reported by Berlau and McGaugh (2006). A BLA role in contextual memory extinction consolidation was demonstrated in our laboratory. Bilateral BLA TTX inactivation, performed after extinction training of this memory task, almost completely impaired extinction (Baldi and Bucherelli, 2010).

### 3.1.1. Neurotransmitter systems

Concerning the different BLA neurotransmitter systems 829 involved in the fear extinction, one of the most investigated is glu-830 tamate and particularly its action at NMDA receptors. Among the 831 first experiments dealing with amygdalar NMDA receptors role in 832 fear extinction there are those by Falls et al. (1992). They employed 833 the fear-potentiated startle paradigm and found that infusion of 834 a NMDA antagonist before extinction training into the BLA blocks 835 extinction of conditioned fear. These results were subsequently 836 replicated not only in fear potentiated startle (Davis, 2002; Lin 837 et al., 2003b) but also using inhibitory avoidance (Myskiw et al., 838

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#### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

### Table 5

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Extinction acquisition: effects of intra-hippocampus pre-extinction training treatments on fear memory extinction. The table shows the studies using local administration of pharmacological or genetic treatments to determine the role of the hippocampus and hippocampal signaling molecules in fear memories extinction acquisition.

Cerebral site	Behavioral paradigm	Treatment	Effect	References
DHC	Auditory fear conditioning	Muscimol	Impairment	Corcoran et al. (2005) and Xue et al. (2014)
VHC	Auditory fear conditioning	Muscimol	Impairment	Sierra-Mercado et al. (2011)
DHC	Auditory fear conditioning	NMDA agonist	Improvement	Ren et al. (2013)
DHC	Contextual fear conditioning	NMDA agonist	Improvement	Bolkan and Lattal (2014)
DHC	Inhibitory avoidance	NMDA antagonist	Impairment	Cammarota et al. (2005), Myskiw et al. (2010) and Szapiro et al. (2003)
DHC	Inhibitory avoidance	CB1 agonist	Improvement	Abush and Akirav (2010)
		CB1 antagonist	Impairment	X
DHC	Inhibitory avoidance	MAPK inhibitor	Impairment	Szapiro et al. (2003)
DHC	Fear-potentiated startle	MAPK inhibitor	No effect	Shen et al. (2011)
DHC	Inhibitory avoidance	PKA inhibitor	Impairment	Myskiw et al. (2010) and Szapiro et al. (2003)
DHC	Inhibitory avoidance	CaMKII	Impairment	Myskiw et al. (2010) and Szapiro et al. (2003)
DHC	Fear-potentiated startle	Ginkgo biloba extract	Improvement	Shen et al. (2011)
DHC	Contextual fear conditioning	SFKs inhibitor	Improvement	Isosaka et al. (2009)
DHC	Contextual fear conditioning	Protein tyrosine phosphatases inhibitor	Impairment	De la Fuente et al. (2011) and Isosaka and Yuasa (2010)
DHC	Inhibitory avoidance	Protein synthesis inhibitor	Impairment	<b>Cammarota et al.</b> (2005) and Vianna et al. (2001, 2003)
DHC	Inhibitory avoidance	mRNA synthesis inhibitor	Impairment	Vianna et al. (2003)
DHC	Contextual fear conditioning	rBDNF	Impairment	Kirtley and Thomas (2010)
		Zif268-ASO	No effect	
DHC	Contextual fear conditioning	NFAT inhibitor	Impairment	De la Fuente et al. (2011)
DHC	Contextual fear conditioning	HDAC inhibitor	Improvement	Lattal et al. (2007)

BDNF, brain-derived neurotrophic factor; CaMKII,  $Ca^{2+}/calmodulin-dependent protein kinase II; CB1, cannabinoid receptor type 1; DHC, dorsal hippocampus; HDAC, histone deacetylase; MAPK, mitogen-activated protein kinase; NFAT, nuclear factor of activated T-cells; NMDA, N-methyl-D-aspartate receptor; PKA, protein kinase A; rBDNF, recombinant BDNF protein; SFKs, Src-family tyrosine kinases; VHC, ventral hippocampus; ZIF268, zinc finger 268; ZIF268-ASO, antisense oligonucleotide targeting ZIF268.$ 

2010), contextual (Kwapis et al., 2014; Laurent et al., 2008; Lee 839 and Kim, 1998) and cued (Holmes et al., 2013; Lee and Kim, 1998; 840 Zimmerman and Maren, 2010) fear conditioning paradigms. These 841 results support the idea that BLA NMDA receptors are involved 842 in acquisition of fear memory extinction. Nevertheless, there is 843 also evidence for amygdalar NMDA receptors role in fear memory 844 extinction consolidation. Intra-BLA infusion of the NMDA antago-845 846 nist immediately following the first of two sessions of extinction 847 produces impairment of inhibitory avoidance and contextual fear extinction (Fiorenza et al., 2012). Further evidence for BLA NMDA 848 receptors role in fear extinction was obtained employing a partial 849 agonist, DCS. DCS facilitates extinction of fear potentiated startle 850 and auditory fear when administered into the BLA before extinc-851 tion training (Davis, 2002; Lee et al., 2006; Mao et al., 2006; Walker 852 et al., 2002) confirming a role of these receptors in extinction 853 acquisition. Moreover, fear extinction facilitation was observed 854 when DCS is injected after extinction training (Akirav et al., 855 2009; Fiorenza et al., 2012; Ledgerwood et al., 2003; Mao et al., 856 2006). This last effect seems to reflect modulation of extinction 857 consolidation (Myskiw et al., 2014). Different results have been 858 obtained using contextual fear paradigm. In fact, intra-BLA DCS 859 administration performed before or immediately after extinction 860 training impairs contextual fear memory extinction. These effects 861 seem to depend on the behavior of the animals during extinction 862 training (Bolkan and Lattal, 2014). 863

The functional role of NMDA receptors seems to depend on their subunit composition (Cull-Candy and Leszkiewicz, 2004). Specifically, the 2B subunit appears to be involved in learning and associated plasticity in several brain sites (Tang et al., 1999; Ge et al., 2007). The NR2B pharmacological manipulation is a relatively selective tool for studying the contribution of NMDA receptormediated plasticity to extinction. The selective inactivation of NMDA receptor containing NR2B subunit in the LA/BLA before extinction training impairs acquisition of conditioned fear extinction to both acoustic CS (Sotres-Bayon et al., 2007) and context (Laurent et al., 2008; Laurent and Westbrook, 2008); on the other hand, the same treatment performed immediately after extinction training has no effect on the extinction of the two fear memory tasks (Laurent and Westbrook, 2008; Sotres-Bayon et al., 2009). These findings suggest that amygdalar NMDA receptors containing 878 NR2B subunit are involved in acquisition, but not consolidation of 879 fear memory extinction. NMDA receptors activation allows calcium 880 influx resulting in an increased intracellular concentration of the 881 ion. However, calcium influx is also associated with L-VGCCs acti-882 vation. The L-VGCCs role in fear extinction is controversial because 883 contradictory results of their involvement were obtained (Schafe, 884 2008) using genetic and pharmacological approaches. A recent 885 work (Davis and Bauer, 2012) employed local infusions into the BLA 886 of L-VGCCs antagonists to test the involvement of these channels in 887 auditory fear extinction. It was found that pre-extinction training 888 L-VGCCs blockade into this neural site induces impairment of long-889 term extinction retention. However, since the animals subjected to 890 this treatment showed extinction acquisition, the results suggest 891 that L-VGCCs are necessary for the fear extinction consolidation. 892

The AMPA receptor is another subtype of glutamate receptors involved in experience-dependent forms of synaptic plasticity (Zushida et al., 2007). Falls et al. (1992) showed that administration of AMPA receptor antagonist into the BLA before extinction training has no effect on subsequent extinction retention of fear potentiated startle. Similar results were obtained in an auditory fear task (Zimmerman and Maren, 2010). Together these results reveal that AMPA receptor in the BLA is not required for fear extinction. However, a recent finding indicates that intra-BLA preextinction training administration of AMPA antagonist impairs contextual fear extinction retention (Kwapis et al., 2014). Moreover, intra-amygdala pre-extinction training injection of an AMPA receptor "potentiator" facilitates contextual fear memory extinction (Zushida et al., 2007). This "potentiator" might exert its effect on fear extinction by promoting AMPA receptors internalization (Kim et al., 2007b; Maren, 2005; Yeh et al., 2006). Recently a synthetic peptide that blocks the internalization of these receptors was employed and it was found that intra-BLA infusion before extinction training impairs auditory fear extinction (Kim et al., 2007b). So the potential role of amygdala AMPA receptors in fear memory extinction requires further investigation.

As recent work has evidenced that glutamate receptors in the CEA are also implicated in the acquisition of fear conditioning, Zimmerman and Maren (2010) investigated the potential role of

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#### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

### 12 Table 6

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Extinction acquisition: effects of intra several cerebral sites pre-extinction training treatments on fear memory extinction. The table shows the studies using local administration of pharmacological or genetic treatments to determine the role of specific cerebral sites and signaling molecules in fear memories extinction acquisition.

Behavioral paradigm	Treatment		References
Auditory fear conditioning	Muscimol	Improvement	Akirav et al. <mark>(2006)</mark>
Auditory fear conditioning	TTX	Impairment	Sierra-Mercado et al. (2006)
	Muscimol	•	Laurent and Westbrook (2008)
0		•	Sierra-Mercado et al. (2011)
		1	Laurent and Westbrook (2009)
0		•	Sierra-Mercado et al. (2011)
			Laurent and Westbrook (2009)
			Cantini and Darter (2010)
Auditory lear conditioning		•	Santini and Porter <mark>(2010</mark> )
			Chang and Maren (2011)
			Burgos-Robles et al. (2007)
	NR2B antagonist		Sotres-Bayon et al. (2009) Laurent and Westbrook (2008)
Contextual fear conditioning	NR2B antagonist	No effect	Laurent and Westbrook (2008)
Contextual fear conditioning	AMPA potentiator	Improvement	Zushida et al. (2007)
Auditory fear conditioning	β-AR antagonist	Impairment	Zushida et al. (2007) Mueller et al. (2008)
Contextual fear conditioning	α-AR antagonist	Impairment	Do Monte et al. (2010)
			Hikind and Maroun (2008)
			Mueller et al. (2010)
			Pfeiffer and Fendt (2006)
		•	Santini et al. (2012)
	ů.		Lin et al. (2009)
	<u> </u>	•	Kubport et al. (2012)
-			Kuhnert et al. (2013) Kuhnert et al. (2013) and Lin et
-			(2009)
Ũ			Do Monte et al. (2013a,b)
Auditory fear conditioning	PKA inhibitor	Impairment	Mueller et al. (2008)
Auditory fear conditioning	CaMKII inhibitor	No effect	Mueller et al. (2008)
Auditory fear conditioning	Protein synthesis inhibitor	Impairment	Mueller et al. (2008) and Santin et al. (2004)
Auditory fear conditioning	mRNA synthesis inhibitor	Impairment	Mueller et al. (2008)
		•	Peters et al (2010)
		1	Peters et al. (2010) Myskiw et al. (2010)
minibitory avoidance			
* * * *			
Inhibitory avoidance			Myskiw et al. (2010)
	PKA inhibitor		
Auditory fear conditioning	D2 antagonist	Impairment	Holtzman-Assif et al. (2010)
Auditory fear conditioning	NMDA antagonist	Impairment	Orsini and Maren (2009)
	AMPA antagonist	Impairment	~
	Protein synthesis inhibitor	No effect	
	MAPK inhibitor	No effect	
Auditory fear conditioning			Padilla-Coreano et al. (2012)
			McNally et al. (2004)
			McNally et al. (2004)
			McNally et al. (2004)
Auditory lear conditioning			McNally et al. (2005)
	•		
	-		
Auditory fear conditioning		•	McNally et al. (2005)
	PKA activator	No effect	. •
	MAPK inhibitor	No effect	
Auditory fear conditioning	Endogenous opioid catabolizing enzymes inhibitor	Improvement	McNally (2005)
Fear-potentiated startle	NMDA antagonist	No effect	Falls et al. (1992)
	Auditory fear conditioning Contextual fear conditioning Auditory fear conditioning Auditory fear conditioning Auditory fear conditioning Auditory fear conditioning Auditory fear conditioning Auditory fear conditioning Contextual fear conditioning Contextual fear conditioning Contextual fear conditioning Auditory fear conditioning	Auditory fear conditioning Contextual fear conditioning Auditory fear conditioning Contextual fear conditioning Auditory fear conditioning Auditory fear conditioning Auditory fear conditioning Auditory fear conditioning Auditory fear conditioning Contextual fear conditioning Auditory fear conditioning Contextual fear conditioning Auditory fear conditioni	Auditory fear conditioning Contextual fear conditioning MuscimolTTXImpairment ImpairmentAuditory fear conditioning Auditory fear conditioning Auditory fear conditioning Auditory fear conditioningMuscimolImpairmentAuditory fear conditioning Auditory fear conditioning Auditory fear conditioningMuscimolNo effectAuditory fear conditioning Auditory fear conditioning 

α-AR, α-adrenergic receptor; AMPA, α-amino-3-hydroxy-5-methyl-4-isoxazole propionic acid receptor; β-AR, β-adrenergic receptor; BDNF, brain-derived neurotrophic factor; CaMKII<sub>4</sub>Ca<sup>2+</sup>/calmodulin-dependent protein kinase II; cAMP<sub>4</sub> cyclic AMP; CB1, cannabinoid receptor type 1; b-ORs<sub>4</sub> opioid receptors subtype b; D1, D2, D4, dopaminergic receptors type 1, 2, 4; dPAG, dorsal periaqueductal gray; IL-mPFC, infralimbic sublegion of the medial prefrontal cortex; κ-ORs, opioid receptors subtype κ; μ-ORs, opioid receptors subtype μ; MAPK, mitogen-activated protein kinase; mPFC, medial prefrontal cortex; NMDA, N-methyl-D-aspartate receptor; NR2B, NMDA receptor subtype 2B; ORs, opioid receptors; PKA, protein kinase A; PL-mPFC, prelimbic subregion of the medial prefrontal cortex; TTX, tetrodotoxin; vIPAG, ventro-lateral periaqueductal gray.

NMDA and AMPA receptors within this amygdaloid nucleus in fear extinction. Neither pre-extinction training NMDA antagonist nor AMPA antagonist injected into the CEA affect auditory fear extinction. Thus, whereas the BLA may have a broader role in acquiring both fear and extinction memories, CEA plays a selective role in fear acquisition.

The literature about the involvement of the metabotropic glutamate receptors (mGluRs) in fear memory extinction is quite limited. The only report, to our knowledge, showed that local infusion of a mGluR1 antagonist into the LA before extinction training impairs the extinction of auditory fear memory (Kim et al., 2007a). Moreover, mGluR1activity seems to be linked specifically to mechanisms

underlying extinction, because intra-LA administration of the same antagonist before fear conditioning has no effect on fear acquisition (Kim et al., 2007a).

 $\gamma$ -Aminobutyric acid (GABA) is considered the major inhibitory 932 neurotransmitter in the mammalian central nervous system. GABA 933 seems to play a complex role in fear extinction and although 934 the results obtained are mixed and sometimes contradictory, this 935 neurotransmitter appears to interfere with the acquisition and 936 consolidation of fear extinction memory. Increasing GABAergic 937 transmission before extinction training disrupts extinction reten-938 tion. For example, Hart et al. (2009, 2010) observed that midazolam 939 (a benzodiazepine) injected intra-BLA before extinction training 940

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#### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

### Table 7

Extinction consolidation: effects of intra-amygdala post-extinction treatments on fear memory extinction. The table shows the studies using local administration of pharmacological or genetic treatments to determine the role of the amygdala and amygdaloid signaling molecules in fear memories extinction consolidation.

Cerebral site	Behavioral paradigm	Treatment	Effect	References
BLA	Auditory fear conditioning	Muscimol	No effect	Sierra-Mercado et al. <mark>(2011)</mark>
BLA	Auditory fear conditioning	Muscimol	Improvement	Akirav et al. (2006)
BLA	Contextual fear conditioning	Muscimol	No effect	Berlau and McGaugh (2006)
BLA	Contextual fear conditioning	Muscimol	Impairment	Laurent and Westbrook (2008)
BLA	Contextual fear conditioning	TTX	Impairment	Baldi and Bucherelli (2010)
BLA	Contextual fear conditioning	GABA antagonist	Improvement	Berlau and McGaugh (2006)
BLA	Fear-potentiated startle	NMDA agonist	Improvement	Mao et al. (2006)
BLA	Inhibitory avoidance	NMDA agonist	Improvement	Fiorenza et al. (2012)
BLA	Cued fear conditioning (CS: light)	NMDA agonist	Improvement	Ledgerwood et al. (2003)
BLA	Contextual fear conditioning	NMDA agonist	Improvement	Akirav et al. (2009) and Fiorenza et al. (2012)
BLA	Contextual fear conditioning	NMDA agonist	Impairment	Bolkan and Lattal (2014)
BLA	Inhibitory avoidance	NMDA antagonist	Impairment	Fiorenza et al. (2012)
BLA	Contextual fear conditioning	NMDA antagonist	Impairment	Fiorenza et al. (2012)
LA	Auditory fear conditioning	NR2B antagonist	No effect	Sotres-Bayon et al. (2009)
BLA	Contextual fear conditioning	NR2B antagonist	No effect	Laurent and Westbrook (2008)
LA	Auditory fear conditioning	β-AR agonist	Impairment	Debiec et al. (2011)
BLA	Contextual fear conditioning	β-AR antagonist	Improvement	Fiorenza et al. (2012)
BLA	Contextual fear conditioning	β-AR antagonist	No effect	Berlau and McGaugh (2006)
BLA	Inhibitory avoidance	β-AR antagonist	Improvement	Fiorenza et al. (2012)
BLA	Contextual fear conditioning	Norepinephrine	No effect	Fiorenza et al. (2012)
BLA	Contextual fear conditioning	Norepinephrine	Improvement	Berlau and McGaugh (2006)
BLA	Inhibitory avoidance	Norepinephrine	Impairment	Fiorenza et al. (2012)
BLA	Contextual fear conditioning	D1 agonist	No effect	Fiorenza et al. (2012)
BLA	Inhibitory avoidance	D1 agonist	No effect	Fiorenza et al. (2012)
BLA	Auditory fear conditioning	D1 antagonist	Impairment	Hikind and Maroun (2008)
BLA	Contextual fear conditioning	D1 antagonist	No effect	Fiorenza et al. (2012)
BLA	Inhibitory avoidance	D1 antagonist	Impairment	Fiorenza et al. (2012)
BLA	Contextual fear conditioning	Muscarinic agonist	Improvement	Boccia et al. (2009)
BLA	Contextual fear conditioning	H2 antagonist	Improvement	Fiorenza et al. (2012)
BLA	Inhibitory avoidance	H2 antagonist	Impairment	Fiorenza et al. (2012)
BLA	Contextual fear conditioning	Histamine N-methyltransferase inhibitor	Improvement	Fiorenza et al. (2012)
BLA	Inhibitory avoidance	Histamine N-methyltransferase inhibitor	Improvement	Fiorenza et al. (2012)
BLA	Auditory fear conditioning	PKA activator	No effect	Tronson et al. (2006)
BLA	Fear-potentiated startle	PI-3 <mark>K i</mark> nhibitor	Impairment	Mao et al. (2006)
BLA	Contextual fear conditioning	PI-3 <mark>K i</mark> nhibitor	Impairment	Kritman and Maroun (2013)
BLA	Auditory fear conditioning	Protein synthesis inhibitor	Impairment	Duvarci et al. (2006)
BLA	Contextual fear conditioning	Protein synthesis inhibitor	Impairment	Mamiya et al. (2009)
BLA	Auditory fear conditioning	FGF2	Improvement	Graham and Richardson (2011)

β-AR, β-adrenergic receptor; BLA, basolateral amygdala; D1, dopaminergic receptor type 1; FGF2, fibroblast growth factor 2; GABA, γ-aminobutyric acid; H2, histaminergic receptor type 2; LA, lateral amygdala; NMDA, N-methyl-b-aspartate receptor; NR2B, NMDA receptor subtype 2B; PI-3K, phosphatidylinositol 3 kinase; PKA, protein kinase A; TTX, tetrodotoxin.

impairs extinction of contextual fear, but spares and facilitates the 941 re-learning of extinction. Thus intra-BLA GABAergic transmission 942 appears to be involved in fear extinction acquisition. However, a 943 study by Akirav et al. (2006) reported findings that seem to con-944 tradict this conclusion. In fact, these authors demonstrated that 945 administration of muscimol (a GABA<sub>A</sub> agonist) into the BLA before 946 an extinction training session does not alter extinction learning of 947 auditory fear conditioning. On the other hand, it was shown that in 948 949 this neural site the levels of gephyrin protein and mRNA and mRNAs of other GABAergic markers (such as GABA-synthesizing enzymes) 950 are significantly increased following extinction training of fear-951 potentiated startle (Chhatwal et al., 2005; Heldt and Ressler, 2007). 952 That is, gephyrin and other GABAergic markers are upregulated 953 after fear extinction training in the BLA suggesting an increased 954 GABAergic transmission (Chhatwal et al., 2005; Heldt and Ressler, 955 2007) 956

GABAergic transmission in the BLA has been implicated in the consolidation of fear extinction as well. Berlau and McGaugh (2006) observed enhanced extinction of contextual fear memory by unilateral intra-BLA infusion of the GABA receptor antagonist bicuculline when performed immediately but not 3 h after extinction training. These results support the idea that intra-BLA GABA antagonists facilitate extinction consolidation. Accordingly, the agonists should impair extinction. Instead, the results by Akirav et al. (2006) showed that administration of muscimol into the BLA after a short extinction session, but not a long one, leads to a significant improvement

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of the auditory fear extinction retention. That is, increasing amyg-967 dalar GABAergic transmission after extinction improves extinction 968 retention. Maybe these unexpected results are due to the method-969 ology used to induce extinction. The authors employed a short 970 extinction session which is more similar to a reactivation session 971 than an extinction one. Therefore they might have disrupted the 972 reconsolidation of the original mnemonic trace rather than facili-973 tated the extinction consolidation. 974

Norepinephrine is implicated in fear learning and memory 975 (Debiec et al., 2011). Noradrenergic involvement in fear extinction 976 was the topic of some recent studies which employed both Pavlo-977 vian fear conditioning and inhibitory avoidance. Results by Berlau 978 and McGaugh (2006) showed that intra-BLA adrenergic activation 979 promotes, whereas adrenergic blockade hinders memory consol-980 idation in fear extinction. Using a contextual fear conditioning 981 paradigm, the authors reported that unilateral intra-BLA infusion 982 of norepinephrine following extinction training enhances extinc-983 tion retention. On the contrary, administration of  $\beta$ -AR antagonist 984 propranolol does not significantly affect extinction retention 985 per se, but it prevents the extinction facilitation by intra-BLA 986 bicuculline (GABAergic antagonist) whether the two compounds 987 are co-administered. Thus, noradrenergic signaling seems to 988 mediate memory modulatory effects of GABA. Different results 989 were obtained by Fiorenza et al. (2012) who considered extinction 990 of two different fear-motivated tasks: contextual fear conditioning 991 and inhibitory avoidance. Intra-BLA post-extinction injection of 992

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E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

### Table 8

Extinction consolidation: effects of intra-hippocampus post-extinction treatments on fear memory extinction. The table shows the studies using local administration of pharmacological or genetic treatments to determine the role of the hippocampus and hippocampal signaling molecules in fear memories extinction consolidation.

Cerebral site	Behavioral paradigm	Treatment	Effect	References
DHC	Contextual fear conditioning	Muscimol	No effect	Berlau and McGaugh <mark>(2006)</mark>
VHC	Auditory fear conditioning	Muscimol	No effect	Sierra-Mercado et al. (2011)
DHC	Inhibitory avoidance	NMDA agonist	Improvement	Fiorenza et al. (2012)
DHC	Contextual fear conditioning	NMDA agonist	Improvement	Bolkan and Lattal (2014) and Fiorenza et al. (2012)
DHC	Inhibitory avoidance	NMDA antagonist	Impairment	Fiorenza et al. (2012) and Szapiro et al. (2003)
DHC	Contextual fear conditioning	NMDA antagonist	Impairment	Pe Carvalho Myskiw et al. (2014) and Fiorenza et al. (2012)
DHC	Contextual fear conditioning	NR2A antagonist NR2B antagonist	Impairment No effect	Leadrebrand et al. (2014)
DHC	Inhibitory avoidance	NR2B agonist <mark>NR</mark> 2B antagonist	No effect No effect	Bonini et al. (2011)
DHC	Contextual fear conditioning	L-VGCC inhibitor	Impairment	De Carvalho Myskiw et al. (2014)
DHC	Inhibitory avoidance	mGluR5 antagonist	No effect	Simonyi et al. (2007)
		mGluR1 antagonist	Impairment	
DHC	Inhibitory avoidance	Norepinephrine	No effect	Fiorenza et al. (2012)
		β-AR antagonist	Impairment	
DHC	Contextual fear conditioning	Norepinephrine	No effect	Fiorenza et al. (2012)
		β-AR antagonist	No effect	
DHC	Contextual fear conditioning	β-AR antagonist	Impairment	Ouyang and Thomas (2005) Fiorenza et al. (2012)
DHC	Inhibitory avoidance	D1agonist	Improvement	Fiorenza et al. (2012)
		D1 antagonist	Impairment	
DHC	Inhibitory avoidance	nAChR agonists nAChR antagonists	Improvement No effect	Pe Aguiar et al. (2013)
DHC	Contextual fear conditioning	D1 agonist D1 antagonist	Improvement Impairment	Fiorenza et al. (2012)
DHC	Contextual fear conditioning	CB1 agonist CB1 antagonist	Improvement Impairment	De Oliveira Alvares et al. (2008)
DHC	Inhibitory avoidance	Histamine N-methyl-transferase inhibitor H2 antagonist	Improvement	Bonini et al. (2011) and Fiorenza et al. (2012)
DUC	× 1 · 1 ·		Impairment	
DHC	Inhibitory avoidance	Histamine, H2 agonist	Improvement	Bonini et al. (2011)
DHC	Inhibitory avoidance	H3 agonist	Impairment	Bonini et al. (2011)
DHC	Inhibitory avoidance	H1 agonist, H1 antagonist, H3 antagonist	No effect	Bonini et al. (2011)
DHC	Contextual fear conditioning	Histamine N-methyl-transferase inhibitor	Improvement Impairment	Fiorenza et ál. (2012)
DHC	Inhibitory avoidance	H2 antagonist GRPR inhibitor	Impairment	Luft at al. (2006)
DHC	Inhibitory avoidance	MAPK inhibitor	Impairment	Luft et al. (2006) Bevilaqua et al. (2007), Bonini et al. (2011),
	-		-	Rossato et al. (2006) and Szapiro et al. (2003)
DHC	Contextual fear conditioning	MAPK inhibitor	Impairment	Fischer et al. (2007) and Huh et al. (2009) Szapiro et al. (2003)
DHC	Inhibitory avoidance	PKA inhibitor AKAPs inhibitor	Impairment	
DHC	Contextual fear conditioning	CaMKII inhibitor	Improvement	Nijholt et al. (2008)
DHC	Inhibitory avoidance		Impairment	Szapiro et al. (2003) Chon et al. (2005)
DHC DHC	Contextual fear conditioning	PI-3 <mark>K</mark> inhibitor Protein synthesis inhibitor	Impairment	Chen et al. $(2005)$ Luft et al. $(2006)$ Power et al. $(2006)$ and
	Inhibitory avoidance		Impairment	Luft et al. (2006), Power et al. (2006) and Vianna et al. (2001)
DHC	Contextual fear conditioning	Protein synthesis inhibitors	Improvement	Fischer et al. (2004)
DHC	Contextual fear conditioning	Protein synthesis inhibitor	Impairment	De Carvalho Myskiw et al. (2014)
Hippocampus	Contextual fear conditioning	Protein synthesis inhibitor	No effect	Mamiya et al. (2009)
DHC	Inhibitory avoidance Contextual fear conditioning	mRNA synthesis inhibitor	No effect	Vianna et al. (2003) Lee et al. (2008)
DHC	•	Proteasome inhibitor	Impairment No effect	De Carvalho Myskiw et al. (2014)
DHC DHC	Contextual fear conditioning Contextual fear conditioning	Proteasome inhibitor CdK5 inhibitor	Improvement	Sananbenesi et al. (2007)
DHC	Contextual fear conditioning	GTPase Rac-1 inhibitor	Improvement	Sananbenesi et al. (2007) Sananbenesi et al. (2007)
DHC	Contextual fear conditioning	PAK-1 inhibitor	Impairment	Sananbenesi et al. (2007) Sananbenesi et al. (2007)
DHC	Contextual fear conditioning	Actin dynamics inhibitors	Impairment	Fischer et al. (2004)
DHC	Inhibitory avoidance	SFKs inhibitor	Impairment	Bevilaqua et al. (2005)
DHC	Contextual fear conditioning	NF-kB inhibitor	Improvement	De la Fuente et al. (2011)
	contextual leaf conditioning		mprovement	

AKAPs, A-kinase anchoring proteins; β-AR, β-adrenergic receptor; CaMKIL, Ca<sup>2+</sup>/calmodulin-dependent protein kinase II; CB1, cannabinoid receptor type 1; Cdk5, cyclindependent kinase 5; D1, dopaminergic receptor type 1; DHC, dorsal hippocampus; GRPR, gastrin-releasing peptide receptor; GTPase Rac<sub>7</sub>1, guanosine triphosphatase Rac-1; H1, H2, H3, histaminergic receptors type 1, 2, 3; L-VGCC, L-type voltage-gated calcium channel; MAPK, mitogen-activated protein kinase; mGluR1, mGluR5, metabotropic glutamate receptors subtype 1, 5; nAChR, nicotinic acetylcholine receptor; NF-κB, nuclear factor κB; NMDA, N-methyl-D-aspartate receptor; NR2A, NR2B, NMDA receptors subtype 2A, 2B; PAK<sub>7</sub>1, p21 activated kinase-1; PI-3K, phosphatidylinositol 3 kinase; PKA, protein kinase A; SFKs, Src-family tyrosine kinases; VHC, ventral hippocampus.

norepinephrine impairs extinction of the inhibitory avoidance, but has no effect on extinction of contextual fear paradigm. Instead, administration of  $\beta$ -AR antagonist timolol in the same neural site enhances extinction of both tasks. Finally, amygdala noradrenergic signaling involvement in fear extinction was shown by Debiec et al. (2011) using an auditory fear conditioning paradigm.  $\beta$ -AR agonist isoproterenol infused into the LA after retrieval of conditioned

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fear impairs its extinction. Although these studies report slightly different results, all seem to indicate an involvement of amygdalar noradrenergic system in the fear extinction consolidation.

Cholinergic activation within the BLA appears to modulate the consolidation of contextual fear memory as well. In fact, intra-BLA infusions of the muscarinic cholinergic agonist oxotremorine performed after each of two extinction sessions cause enhanced

#### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

### Table 9

Extinction consolidation: effects of intra several cerebral sites post-extinction treatments on fear memory extinction. The table shows the studies using local administration of pharmacological or genetic treatments to determine the role of specific cerebral sites and signaling molecules in fear memories extinction consolidation.

Cerebral site	Behavioral paradigm	Treatment	Effect	References
mPFC	Auditory fear conditioning	Muscimol	No effect	Akirav et al. <mark>(2006</mark> ) Sierra-Mercado et al. <mark>(2006)</mark>
mPFC	Auditory fear conditioning	TTX	No effect	Sierra-Mercado et al. (2006)
mPFC	Contextual fear conditioning	Muscimol	Impairment	Laurent and Westbrook (2008)
IL-mPFC	Auditory fear conditioning	Muscimol	No effect	Sierra-Mercado et al. (2011)
IL-mPFC	Contextual fear conditioning	Muscimol	Impairment	Laurent and Westbrook (2009)
IL-mPFC	Auditory fear conditioning	M-type K <sup>+</sup> channels antagonist	No effect	Santini and Porter (2010)
IL-mPFC	Auditory fear conditioning	GABA antagonist	Improvement	Chang and Maren (2011)
IL-mPFC	Contextual fear conditioning	GABA antagonist	Improvement	Thompson et al. <mark>(2010</mark> )
PL-mPFC	Contextual fear conditioning	GABA antagonist	No effect	Thompson et al. (2010)
mPFC	Contextual fear conditioning	NMDA agonist	Improvement	Fiorenza et al. (2012)
mPFC	Inhibitory avoidance	NMDA agonist	Improvement	Fiorenza et al. (2012)
mPFC	Auditory fear conditioning	NMDA antagonist	Impairment	Burgos-Robles et al. (2007) and Holmes et al. (2012)
mPFC	Contextual fear conditioning	NMDA antagonist	Impairment	Fiorenza et al. (2012)
mPFC	Inhibitory avoidance	NMDA antagonist	Impairment	Fiorenza et al. (2012)
mPFC	Auditory fear conditioning	NR2B antagonist	Impairment	Sotres-Bayon et al. (2009)
mPFC	Contextual fear conditioning	NR2B antagonist	Impairment	Laurent and Westbrook (2008)
IL-mPFC	Auditory fear conditioning	β-AR antagonist	No effect	Mueller et al. (2008)
mPFC	Contextual fear conditioning	β-AR antagonist	Improvement	Fiorenza et al. (2012)
mPFC	Inhibitory avoidance	β-AR antagonist	Impairment	Fiorenza et al. (2012)
mPFC	Contextual fear conditioning	Norepinephrine	Impairment	Fiorenza et al. (2012)
mPFC	Inhibitory avoidance	Norepinephrine	No effect	Fiorenza et al. (2012)
mPFC	Contextual fear conditioning	D1 agonist	No effect	Fiorenza et al. (2012)
mPFC	Inhibitory avoidance	D1 agonist	No effect	Fiorenza et al. (2012)
IL-mPFC	Auditory fear conditioning	D1 antagonist	Impairment	Hikind and Maroun (2008)
mPFC	Contextual fear conditioning	D1 antagonist	No effect	Fiorenza et al. (2012)
mPFC	Inhibitory avoidance	D1 antagonist	Impairment	Fiorenza et al. (2012)
mPFC	Contextual fear conditioning	H2 antagonist	Impairment	Fiorenza et al. (2012)
mPFC	Inhibitory avoidance	H2 antagonist	Impairment	Fiorenza et al. (2012)
mPFC	Contextual fear conditioning	Histamine N-methyltransferase inhibitor	Improvement	Fiorenza et al. (2012)
mPFC	Inhibitory avoidance	Histamine N-methyltransferase inhibitor	No effect	Fiorenza et al. (2012)
mPFC	Auditory fear conditioning	MAPK inhibitor	Impairment	Hugues et al. (2004)
IL-mPFC	Contextual fear conditioning	PI-3 <mark>K</mark> inhibitor	Impairment	Kritman and Maroun (2013)
mPFC	Contextual fear conditioning	Protein synthesis inhibitor	Impairment	Mamiya et al. (2009)
ENT	Inhibitory avoidance	Protein synthesis inhibitor	Impairment	Bevilaqua et al. <mark>(2006)</mark>
		NMDA antagonist	Impairment	~
		CaMKII antagonist	Impairment	
		MAPK inhibitor	No effect	
ENT	Contextual fear conditioning	TTX	Impairment	Baldi and Bucherelli (2014)
Nucleus basalis magnocellularis	Contextual fear conditioning	TTX	No effect	Baldi and Bucherelli (2010)
<mark>S</mark> ubstantia nigra	Contextual fear conditioning	TTX	No effect	Baldi and Bucherelli (2010)

β-AR, β-adrenergic receptor; CaMKIL Ca<sup>2+</sup>/calmodulin-dependent protein kinase II; D1, dopaminergic receptor type 1; ENT, entorhinal cortex; GABA, γ-aminobutyric acid; H2, histaminergic receptor type 2; IL-mPFC, infralimbic subregion of the medial prefrontal cortex; MAPK, mitogen-activated protein kinase; mPFC, medial prefrontal cortex; NMDA, N-methyl-D-aspartate receptor; NR2B, NMDA receptor subtype 2B; PI-3K, phosphatidylinositol 3 kinase; PL-mPFC, prelimbic subregion of the medial prefrontal cortex; TTX, tetrodotoxin.

fear extinction (Boccia et al., 2009). The effect is time-dependent because the same treatment administered 180 min after extinction does not affect extinction memory. This provides evidence that oxotremorine facilitates consolidation of extinction.

Previous results have shown that the histaminergic system modulates consolidation of some fear memories (Cangioli et al., 2002; Passani et al., 2001). Recent researches have reported that this system modulates fear extinction as well. Post-extinction infusions into the BLA of a histaminergic H2 receptor antagonist hinder extinction retention of both contextual fear conditioning and inhibitory avoidance (Fiorenza et al., 2012). On the contrary, intra-BLA administration of a histamine-N-methyltransferase inhibitor after extinction training enhances extinction memory retention of both tasks (Fiorenza et al., 2012). Thus, the histaminergic system modulates through H2 receptor extinction consolidation of fear memory (Myskiw et al., 2014).

The dopaminergic system modulates learning during fear extinction (Abraham et al., 2014). This evidence comes from studies employing either systemic administration of dopaminergic receptors agonists or antagonists, or mice lacking these receptors (Hikind and Maroun, 2008). The dopaminergic receptors are highly expressed in the amygdala; specifically, the BLA expresses D1 receptors, whereas the CEA mainly expresses D2 receptors. It

was shown that microinfusions of a D1 antagonist in the BLA before 1030 an extinction session of an auditory fear task cause impairment 1031 in extinction acquisition. However, the same treatment performed 1032 immediately after the extinction session has no effect on the subse-1033 quent extinction retention (Hikind and Maroun, 2008). Thus, these 1034 results are consistent with the idea that fear extinction acqui-1035 sition, but not consolidation, depends on the BLA D1 receptors. 1036 Further analysis of fear extinction consolidation was performed 1037 by Fiorenza et al. (2012) employing the contextual fear condition-1038 ing and inhibitory avoidance paradigms. These authors injected 1039 intra-BLA D1 agonist or antagonist after the first of two sessions of 1040 extinction in each task to influence extinction consolidation. While 1041 D1 agonist had no effect on the extinction of the two tasks, the D1 1042 antagonist impaired extinction consolidation of inhibitory avoid-1043 ance, but not contextual fear conditioning. Together these results 1044 implicate amygdalar dopaminergic signaling as a critical modula-1045 tory component in fear extinction. 1046

Recently, the endocannabinoid system has emerged as an important system in the regulation of extinction learning. Mutant mice lacking the gene for the CB1 receptor  $(CB1^{-/-})$  acquire and retain an auditory fear conditioning task, but show an impairment of extinction acquisition and retention (Marsicano et al., 2002). Moreover, the wild type mice  $(CB1^{+/+})$  exhibit an impaired 1052

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### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

extinction of fear memory when administered a CB1 antagonist before, but not after, extinction training. In wild type mice an increased expression of two endocannabinoids in the BLA was reported following extinction training. Subsequently, it was found that a CB1 antagonist injected into the BLA either before or after extinction training impaired extinction of inhibitory avoidance (Ganon-Elazar and Akirav, 2009). Thus, the CB1 receptor in this neural site is crucially involved in extinction of this memory task. However, the administration in the BLA of a CB1/CB2 agonist had no effect on inhibitory avoidance extinction (Ganon-Elazar and Akirav, 2009). Similar results were obtained using an inhibitor of cannabinoid reuptake and enzymatic degradation (Ganon-Elazar and Akirav, 2009). The amygdalar CB1 receptors involvement in fear-potentiated startle appears to be different. In fact, in this memory task neither CB1 antagonist nor agonist administration into the BLA affects extinction retention (Kuhnert et al., 2013).

Neuropeptide S (NPS) has anxiolytic-like effects and seems to be involved in fear memory extinction (Jungling et al., 2008). Pre-extinction training infusion of NPS within the BLA facilitated auditory fear extinction retention; in addition, the intra-BLA administration of a NPS receptor antagonist induced a significant impairment of extinction learning (Jungling et al., 2008).

Evidence has emerged correlating glucocorticoids release to the fear extinction memory (Rodrigues et al., 2009). The BLA is one of the neural sites containing GRs and thus it may represent a site where extinction memory is modulated. An increase in plasma corticosterone levels was observed following extinction training of the fear potentiated startle in rat suggesting an involvement of this hormone in the plasticity related to the extinction. Consistent with this, systemic administration of a glucocorticoid agonist performed pre- or post-extinction training facilitated fear extinction. Probably this effect is mediated at the level of the amygdala because intra-BLA infusion of a glucocorticoid agonist before extinction training improves extinction (Yang et al., 2006). Moreover, administration of glucocorticoid antagonist either systemically or intra-BLA before extinction training impairs extinction of conditioned fear (Yang et al., 2006). Thus, corticosterone contributes to fear extinction acting at amygdaloid GRs.

### 3.1.2. Protein kinases and phosphatases

Several intracellular signaling pathways have been implicated in fear extinction. MAPK is one of the second messengers activated by increased intracellular calcium concentration following extinction training. It was demonstrated that intra-BLA administration of MAPK inhibitors before extinction training impaired extinction retention in both fear-potentiated startle (Davis, 2002; Lin et al., 2003b; Lu et al., 2001) and auditory fear paradigm (Herry et al., 2006). Moreover, a few studies indicate that phosphorylated MAPK is upregulated into this neural site following extinction training (Cannich et al., 2004; Davis and Bauer, 2012; Kwapis et al., 2014; Yang and Lu. 2005).

Little evidence is available for a role of PKA in fear extinction. Tronson et al. (2006) found that intra-BLA infusions of a specific PKA activator after each of four daily extinction training sessions have no effect on auditory fear extinction. On the other hand, Myskiw et al. (2010) reported that administration of a PKA inhibitor within this neural site prior to the first of several extinction sessions hinders inhibitory avoidance extinction. However, the Authors did not report any change in the amygdaloid phosphorylated PKA levels and suggested that "probably the basal levels of PKA activity in BLA are necessary and sufficient for extinction processes to develop in this task". This conclusion, however, was not confirmed by experiments in which transgenic mice with reduced PKA activity in forebrain neurons were used. In fact, in these mice the reduction of PKA activity facilitates extinction retention of contextual fear (Isiegas et al., 2006). Myskiw et al. (2010) also investigated the amygdaloid CaMKII role in inhibitory avoidance extinction. Using the same protocol of inhibitory avoidance they showed that intra-BLA pre-extinction CaMKII inhibition impairs extinction of this task. This effect is correlated with an increase in phosphorylated CaMKII levels 90 and 180 min after testing.

It has been suggested that fear extinction is correlated with amygdaloid PI-3K activation. Some studies reported that intra-BLA administration of PI-3K inhibitors performed before extinction 1125 training impairs extinction of both fear-potentiated startle (Lin 1126 et al., 2003b; Mao et al., 2006) and contextual fear conditioning 1127 (Kritman and Maroun, 2013). On the other hand, Yang and Lu (2005) 1128 showed that intra-BLA blockade reduces fear-potentiated startle 1129 extinction facilitation induced by systemic, pre-extinction train-1130 ing injection of DCS. PI-3K activity is often evaluated employing 1131 pAKT levels as an indirect measure, however inconsistent results 1132 were obtained. Indeed, following extinction amygdaloid pAKT may 1133 increase (Yang and Lu, 2005), decrease (Lin et al., 2003a) or remain 1134 unchanged (Cannich et al., 2004). 1135

These results suggest that several kinases in the BLA are involved 1136 in fear extinction. Many of these kinases are dephosphorylated, 1137 that is inactivated, by the protein phosphatase calcineurin. This 1138 phosphatase in the BLA has been implicated in fear extinction. Several works reported that extinction is associated with enhanced 1140 BLA calcineurin levels and enzymatic activity (and consequent 1141 reduced phosphorylation of MAPK and AKT) (Cannich et al., 2004; 1142 Lin et al., 2003a, 2003b). Furthermore, pre-extinction intra-BLA 1143 administration of inhibitors of this phosphatase blocks extinction of 1144 fear-potentiated startle (Lin et al., 2003a). These results suggested 1145 that fear extinction may involve the reversal of acquisition-related 1146 plasticity through upregulation of calcineurin, hence weakening 1147 the original fear memory (Lin et al., 2003a, 2003b).

### 3.1.3. Gene expression and protein synthesis

As recalled previously, the protein kinases activate some tran-1150 scription factors, such as CREB which plays a critical role in fear 1151 memory consolidation (Kida et al., 2002). In fear extinction the role 1152 of CREB is contradictory. Some studies reported increased CREB 1153 phosphorylation after fear extinction (Hall et al., 2001; Mamiya 1154 et al., 2009) but others demonstrated a decreased CREB activity 1155 after extinction (Izumi et al., 2008; Lin et al., 2003b). Recently, a 1156 study by Tronson et al. (2012) attempted to clarify the role of amyg-1157 dalar CREB in this phase of fear memory. Using an auditory fear 1158 conditioning task and intra-BLA CREB viral vectors injection, the 1159 authors showed that extinction is not affected by either disruption 1160 or overexpression of CREB. These results, therefore, seem to sup-1161 port the hypothesis that CREB activity in the BLA is not required for 1162 fear extinction. On the contrary, transcription factor BDNF appears 1163 to be implicated in fear memory extinction. In fact, it was found 1164 that BDNF mRNA expression within the BLA is increased in a time-1165 dependent manner following fear-potentiated startle extinction 1166 (Chhatwal et al., 2006). As BDNF acts on tyrosine kinase B (TrkB) 1167 receptor, the involvement of this receptor in extinction was inves-1168 tigated as well. It was reported that intra-BLA infusion of TrkB 1169 lentiviral vector before extinction training induces a retention, but 1170 not acquisition, deficit of extinction suggesting a BDNF role in fear 1171 memory extinction consolidation (Chhatwal et al., 2006). 1172

BDNF is not the only neurotrophic factor implicated in fear 1173 extinction. A recent work suggested that the fibroblastic growth 1174 factor 2 (FGF2) might be an attractive candidate for enhancing 1175 the learning processes underlying fear extinction (Graham and 1176 Richardson, 2011). Adult FGF2-treated rats exhibit facilitated audi-1177 tory fear extinction consolidation when this neurotrophic factor is 1178 administered within the BLA immediately after extinction training. 1179 These animals also show attenuated renewal and reinstatement of 1180 fear. Therefore, FGF2 appears to be a powerful modulator of fear 118 extinction. 1182

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Fear extinction is partly explained as a new learning which involves the formation of a second, inhibitory association. This new association has opposite effects than the excitatory one, as the CS presentation no longer predicts the US and no fear is expressed. As a new learning, fear extinction requires new protein synthesis. Many studies using intra-BLA pre- or post-extinction training infusion of the protein synthesis inhibitor anisomycin showed an extinction impairment considering several conditioned fear paradigms, such as fear-potentiated startle (Lin et al., 2003b; Yang and Lu, 2005), auditory (Duvarci et al., 2006) and contextual fear conditioning (Mamiya et al., 2009). Probably protein synthesis that occurs during extinction depends on the transcription of new RNA. In this regard, the experiments employing transcription inhibitors showed conflicting results. Lin et al. (2003b) showed that intra-BLA administration of actinomycin D before extinction training has no effect on fear-potentiated startle extinction. It should be underlined that these authors measured extinction retention 20 min after extinction training. In general, short-term memory extinction (which is independent on transcription) is evaluated at a brief interval. Using a fear-potentiated startle paradigm, Yang and Lu (2005) found that extinction facilitation induced by intra-BLA pre-extinction training administration of DCS is blocked by preinjection of actinomycin D into the BLA. These results seem to support the idea that fear memory extinction requires new mRNA synthesis in this neural structure.

Several post-translational modification occur during memory formation. The neural cell adhesion molecule (NCAM), a glycopro-1209 tein of the immunoglobulin superfamily, participates in changes. 1210 For example, polysialylated NCAM (PSA-NCAM) is upregulated 1211 within the amygdala (BLA and CEA) and hippocampus 24 h after 1212 training of auditory fear conditioning and spatial memory, respec-1213 tively (Markram et al., 2007; Venero et al., 2006). In the amygdala 1214 this upregulation is not necessary for the acquisition, consolida-1215 tion and recall of fear memories, but it is involved in extinction 1216 of these memories. In fact, pre- and post-training cleavage of PSA 1217 from NCAM induced by enzyme endoneuraminidase N (endoN) 1218 does not interfere with the acquisition or the consolidation of 1219 auditory and contextual fear memories. On the contrary, pre-1220 extinction intra-BLA administration of endoN improves tone fear 1221 memory extinction (Markram et al., 2007). Thus, PSA-NCAM might 1222 be considered a molecular process that plays different roles in the 1223 acquisition and extinction of auditory fear memories, as it occurs 1224 with other mechanisms (Lin et al., 2003a). 1225

### 1226 3.2. Hippocampus

Fear extinction is a context-dependent process; a contex-1227 tual change causes a renewal of extinguished conditioned fear 1228 responses that are again expressed (Herry et al., 2010; Myers and 1229 Davis, 2007). The hippocampus plays a critical role in the formation 1230 of contextual representations (Fanselow, 2000; Kim and Fanselow, 1231 1992; Phillips and LeDoux, 1992), therefore many studies inves-1232 tigated its role in the contextual modulation of fear extinction 1233 (Bouton et al., 2006; Ji and Maren, 2007). Nevertheless, the hip-1234 pocampus appears directly implicated in extinction acquisition, but 1235 not consolidation, of some type of fear memory (Tables 5 and 8). 1236 Pre-extinction training muscimol-induced inactivation of dorsal 1237 hippocampus (DHC) attenuates the extinction acquisition of con-1238 ditioned freezing response to an acoustic CS (Corcoran et al., 2005; 1239 Xue et al., 2014). Instead, unilateral infusion of muscimol into the 1240 DHC immediately after contextual fear extinction training does 1241 not affect this conditioned response (Berlau and McGaugh, 2006). 1242 Whereas pre-extinction muscimol infusion within ventral hip-1243 pocampus (VHC) impairs auditory fear extinction retention, the 1244 1245 same treatment performed immediately after extinction training 1246 has no effect on extinction memory (Sierra-Mercado et al., 2011).

Thus, the activity necessary for fear extinction processing in the hippocampus seems to occur during extinction training. 1248

### 3.2.1. Neurotransmitter systems

In the hippocampus, as in the amygdala, various neurotrans-1250 mitter systems appear critical for fear extinction, depending on the 1251 nature of the mnemonic task. Hippocampal glutamatergic neuro-1252 transmission is involved in extinction of fear memory, although 1253 the ionotropic and metabotropic receptors of glutamate appear to 1254 be implicated to a different extent. Hippocampal NMDA receptor 1255 activation is necessary for the transduction cascade that mediates 1256 the plasticity underlying fear memory extinction. Pre-(Cammarota 1257 et al., 2005; Myskiw et al., 2010; Szapiro et al., 2003) or post-1258 extinction (De Carvalho Myskiw et al., 2014; Fiorenza et al., 2012; 1259 Szapiro et al., 2003) DHC infusion of NMDA antagonists impairs 1260 inhibitory avoidance and contextual fear long-term extinction. 1261 This effect does not seem to be mediated by the NR2B subunit 1262 (Bonini et al., 2011; Leadrebrand et al., 2014). However, hip-1263 pocampal NR2A activity seems to be required for contextual fear 1264 extinction because its blockade performed each day after the 1265 extinction session impairs extinction (Leadrebrand et al., 2014). 1266 Also, intra-hippocampus infusion of DCS before extinction facil-1267 itates acquisition and retrieval of auditory and contextual fear 1268 extinction memory (Bolkan and Lattal, 2014; Ren et al., 2013). Sim-1269 ilarly, post-extinction injections of DCS or p-serine in the same 1270 neural site enhance extinction consolidation of inhibitory avoid-1271 ance and contextual fear task (Bolkan and Lattal, 2014; Fiorenza 1272 et al 2012) 1273

The role of hippocampal mGluRs is less clear. It was shown that mGluR5 knock-out mice exhibit a complete deficit in auditory and contextual fear extinction (Xu et al., 2009). However, this receptor is not involved in extinction of inhibitory avoidance. In fact, mGluR5 antagonist injected intra-DHC after the first extinction session has no effect on subsequent extinction retention of this task (Simonyi et al., 2007). On the contrary, mGluR1 blockade induces a significant impairment of inhibitory avoidance extinction (Simonyi et al., 2007). Thus, extinction of Pavlovian fear conditioning and inhibitory avoidance seems to involve different mGluR subtypes.

Also hippocampal L-VGCCs are crucial for contextual fear extinction. Intra-DHC infusion of nifedipine given after extinction session impairs consolidation of this memory phase (De Carvalho Myskiw et al., 2014). This effect is blocked by the co-administration of the proteasome inhibitor  $\beta$ -lac suggesting that L-VGCCs action depends on concomitant synaptic protein turnover (De Carvalho Myskiw et al., 2014).

Immediately post-extinction training administration of norepinephrine within the DHC has no effect on the extinction of either contextual freezing or inhibitory avoidance (Fiorenza et al., 2012). However, hippocampal  $\beta$ -ARs blockade results in the impairment of extinction consolidation of inhibitory avoidance, but not of contextual freezing response (Fiorenza et al., 2012). In the latter paradigm Ouyang and Thomas (2005) reported that  $\beta$ -adrenergic antagonism within DHC blocks extinction when the treatment is performed 3 h after extinction training, but not when performed before extinction training.

There is little evidence on hippocampal dopaminergic trans-1301 mission involvement in fear extinction. The only study, to our 1302 knowledge, showed that intra-hippocampus administration of D1 1303 receptor agonists or antagonists after extinction training enhances 1304 or impairs, respectively, extinction of both contextual fear and 1305 inhibitory avoidance (Fiorenza et al., 2012). Just as there are 1306 few studies on the role of hippocampal cholinergic receptors. 1307 De Aguiar et al. (2013) found that nicotine and its metabolite 1308 cotinine (nicotinic acetylcholine receptors (nAChRs) agonists) 1309 enhance extinction of inhibitory avoidance when they are injected 1310

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#### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

intra-DHC after the first extinction session. Instead, nAChRs antagonists do not significantly interfere with this mnemonic process.

As reported for the BLA, hippocampal cannabinoid system is involved in fear extinction as well. In fact, infusion into the CA1 of CB1 antagonist before the first extinction session impairs the extinction of inhibitory avoidance, while CB1 agonist or an inhibitor of endocannabinoid reuptake facilitate it (Abush and Akirav, 2010). Similar findings were obtained using a contextual fear paradigm and post-extinction training injections: CB1 antagonists block whereas CB1 agonists enhance the extinction of this memory task (De Oliveira Alvares et al., 2008).

The hippocampal activity is also modulated by histamine. Recently, it was reported that histaminergic system of this neural site is involved in fear extinction consolidation. In particular, it seems that histamine facilitates fear extinction consolidation through a mechanism involving hippocampal H2 receptors. Animals subjected to post-extinction administration of histamine or histamine N-methyl-transferase inhibitor or H2 agonists exhibit enhanced extinction of inhibitory avoidance and contextual freezing response (Bonini et al., 2011; Fiorenza et al., 2012). Opposite effects were observed following injections of H2 antagonist and H3 agonist; moreover, H2 antagonism blocks histamine-induced facilitation (Bonini et al., 2011; Fiorenza et al., 2012). On the contrary, H1 agonists and antagonists or H3 antagonists have no effect on fear extinction (Bonini et al., 2011).

Hippocampal molecular mechanisms mediating extinction of inhibitory avoidance include activation of the gastrin-releasing peptide receptor (GRPR) because its inhibition immediately after the first extinction session blocks this memory phase (Luft et al., 2006).

### 3.2.2. Protein kinases

The many neurotransmitter systems mentioned above activate signaling pathways such as MAPK, PKA and CaMKII. It was reported that inhibitory avoidance and contextual fear extinction is blocked by intra-DHC MAPK (Bevilagua et al., 2007; Bonini et al., 2011; Fischer et al., 2007; Huh et al., 2009; Rossato et al., 2006; Szapiro et al., 2003), PKA (Myskiw et al., 2010; Szapiro et al., 2003), and CaMKII (Myskiw et al., 2010; Szapiro et al., 2003) inhibitors regardless of whether they were given before or after extinction sessions. However, intra-DHC pre-extinction administration of a MAPK (in particular ERK1/2) inhibitor does not affect extinction of fear potentiated startle (Shen et al., 2011). This might be due to the fact that diverse memory tasks use different MAPK subfamilies to produce extinction. It was reported that the subfamily ERK1/2 is not involved in the facilitation effect of Ginkgo biloba extract on fear extinction. In fact, intra-hippocampal infusion of this extract given prior to a single extinction session facilitates conditioned fear extinction as measured by fear-potentiated startle but this effect is only partially attenuated by ERK1/2 inhibitor injection and does not reach a significant level (Shen et al., 2011).

The role of ERK in contextual fear extinction was confirmed using transgenic animals. Rap2V12 transgenic mice express constitutively active Rap 2 (a Rap GTPase of the Ras family) in postnatal forebrain including the hippocampus. These animals exhibit normal conditioned fear acquisition, but impaired contextual fear extinction associated with decreased hippocampal ERK activity after the second and third extinction sessions compared to wildtype controls. This effect may be ascribed to active Rap2 repressing ERK signaling (Ryu et al., 2008). The hippocampus appears to be a major site of Rap2 action because Rap2V12 mice show normal extinction of auditory fear memory and normal amygdaloid and cortical ERK activation (Ryu et al., 2008).

ERK-1 knockout mice are characterized by stimulus-dependent overactivation of the ERK2 isoform and therefore have been used to study the selective role of this ERK isoform. These mice exhibit enhanced contextual fear extinction accompanied by faster and stronger activation of ERK2 than their wild-type littermates (Tronson et al., 2008). According to the authors, these findings support the idea that ERK2 compensates for the lack of ERK1 and shows stronger biological activity in the absence of ERK1 (Tronson et al., 2008). Moreover, intra-hippocampal infusion of a MEK inhibitor after each daily extinction session reduces ERK phosphorylation in both ERK-1 deficient mice and wild-type mice, whereas intra-DHC administration of a PKA or PKC inhibitor does not affect pERK level. Thus, these results indicate a key role of MEK, but not PKA nor PKC, in hippocampal ERK regulation during extinction (Tronson et al., 2008).

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Transgenic mice have also been employed to further clarify the PKA role in contextual fear memory extinction. TetO-R(AB) transgenic mice with reduced PKA activity in the forebrain exhibit facilitated contextual fear extinction retention compared with wild-type controls (Isiegas et al., 2006) suggesting an inhibitory role of PKA in this mnemonic phase. PKA signaling is partly controlled by association of the enzyme with A-kinase anchoring proteins (AKAPs). Nijholt et al. (2008) found that inhibition of hippocampal PKA anchoring AKAPs after each extinction session facilitates contextual fear memory extinction, confirming that PKA activity inhibits extinction process.

Extinction process also seems to require downregulation of PKC signaling. In fact, intra-DHC post-extinction administration of PKC inhibitor facilitates conditioned contextual freezing extinction. However, this treatment does not affect ERK activity suggesting that PKC suppresses fear extinction through an ERK-independent mechanism (Tronson et al., 2008).

It was recently shown that contextual fear extinction involves 1405 hippocampal PI-3K (Chen et al., 2005) and cyclin-dependent kinase 1406 5 (CdK5) (Sananbenesi et al., 2007) pathways. Animals infused 1407 with PI-3K inhibitor into the DHC immediately after repeated tests 1408 do not exhibit decrease in the contextual conditioned freezing 1409 (Chen et al., 2005). On the other hand, inhibition of hippocam-1410 pal CdK5 facilitates extinction of this fear response (Sananbenesi 1411 et al., 2007). Similar results were obtained by intra-hippocampal 1412 injections of upstream regulator GTPase Rac-1 (guanosine triphos-1413 phatase Rac-1) inhibitor. Thus, Rac-1 and CdK5 activity seems to 1414 inhibit contextual fear extinction. Furthermore, it was reported 1415 that downstream target PAK-1 (p21 activated kinase-1) is also 1416 involved in this memory phase because its inhibition within DHC 1417 after extinction training impairs contextual conditioned freezing 1418 response extinction (Sananbenesi et al., 2007). Therefore, the hip-1419 pocampal Rac-1/CdK5/PAK-1 pathway is important for contextual 1420 fear extinction. This pathway appears to affect the dynamics of the 1421 actin cytoskeleton, whose rearrangement in the DHC is required 1422 for extinction process. Indeed, post-extinction training administra-1423 tion of actin dynamics inhibitors impairs contextual conditioned 1424 freezing extinction (Fischer et al., 2004). 1425

Several members of the Src-family tyrosine kinases (SFKs) in the 1426 hippocampus are involved in extinction of fear-motivated memo-1427 ries as well. Using the inhibitory avoidance paradigm Bevilaqua 1428 et al. (2005) found that hippocampal infusion of a specific SFKs 1429 inhibitor performed immediately after the first of four extinc-1430 tion sessions blocks memory extinction and they suggested that 1431 SFKs play a role in consolidation of inhibitory avoidance extinc-1432 tion. Other authors have reported that pre-extinction inhibition 1433 of this kinases family within DHC facilitates extinction of con-1434 textual conditioned freezing; the facilitated extinction is related 1435 to downregulation of hippocampal Fyn activity, a member of 1436 SFKs (Isosaka et al., 2009). Moreover, hippocampal pre-extinction 1437 training administration of protein tyrosine phosphatases inhibitor 1438 impairs extinction of contextual fear memory (Isosaka and Yuasa, 1439 2010). The authors speculated that during extinction training an 1440 increased activity of the protein tyrosine phosphatases might 1441

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occur and these phosphatases should directly or indirectly dephosphorylate Fyn. On the other hand, hippocampal phosphatases
involvement in contextual fear memory extinction was reported
also by De la Fuente et al. (2011), Intra-DHC inhibition of calcineurin
performed before extinction training is related to high levels of
contextual freezing response indicating impaired extinction.

### <sup>1448</sup> 3.2.3. *Gene expression and protein synthesis*

As previously stated, there is evidence supporting the role of 1449 gene transcription during contextual long-term memory extinc-1450 tion in amygdala and prefrontal cortex (Mamiya et al., 2009). 1451 Surprisingly, it was demonstrated that CREB is not activated in 1452 the hippocampus during contextual fear extinction suggesting 1453 that gene expression in this neural site may not be involved 1454 (Mamiya et al., 2009; Tronson et al., 2008). Nevertheless, Kirtley 1455 and Thomas (2010) showed that intra-DHC infusion of recombi-1456 nant BDNF protein before contextual extinction training impairs 1457 consolidation, but not acquisition, of extinction, whereas antisense 1458 oligodeoxynucleotide targeting Zif268 injection does not affect 1459 extinction process. Moreover, Peters et al. (2010) reported that rats 1460 failing to learn extinction of auditory fear show reduced BDNF in hippocampal inputs to the IL-mPFC and enhancing BDNF in this pathway allows extinction of this memory task. These data are consistent with studies in which genetic knockdown of hippocampal BDNF impairs conditioned fear extinction as measured both with 1465 fear potentiated startle and contextual freezing (Heldt et al., 2007). 1466

**De la Fuente et al. (2011) studied the role of two related** transcription factors, NF- $\kappa$ B and NFAT (nuclear factor of activated Tcells), in the hippocampus in extinction memory formation. These transcription factors have opposite roles; in fact, post-extinction training NF- $\kappa$ B inhibition within DHC enhances contextual fear extinction, whereas NFAT blockade performed before extinction training impairs this memorization phase. The authors proposed that the activation or inhibition of these two transcription factors should be regulated by calcineurin phosphatase; during extinction calcineurin might block NF- $\kappa$ B activation and activate NFAT (De la Fuente et al., 2011).

Recent evidence demonstrated that immediate early genes involved in fear extinction are not the same implicated in fear acquisition. For example, during contextual fear conditioning c-Fos and JunB are upregulated (Huff et al., 2006; Strekalova et al., 2003), whereas their expression decreases during subsequent exposures to the same context (Guedea et al., 2011; Tronson et al., 2009) suggesting that they are not activated by extinction. On the contrary, JunD is activated by contextual fear extinction but is not affected by fear acquisition (Guedea et al., 2011). Thus, the learning processes underlying acquisition and extinction of fear are partially different at a molecular level.

Finally, it was shown that extinction of contextual fear conditioning may be modulated by manipulating HATs and HDACs activity. In particular, an HDAC inhibitor (which blocks histone deacetylases activity and increases histone acetylation) given in the hippocampus immediately before extinction training enhances retention of conditioned freezing extinction (Lattal et al., 2007). This effect is due to enhancement of consolidation as the HDAC inhibitor begins to affect histone acetylation about 30 min after administration.

The role of hippocampal protein synthesis in fear extinction has been widely studied. Using an inhibitory avoidance paradigm it was found that intra-hippocampal (dorsal CA1 region) administration of the protein synthesis inhibitor anisomycin, performed either before (Cammarota et al., 2005; Vianna et al., 2001, 2003) or immediately after (Luft et al., 2006; Power et al., 2006; Vianna et al., 2001) the first extinction session, blocks extinction of this fear memory task. Instead, the treatment is ineffective when given 1 or 3 h after the first extinction session (Vianna et al., 2003). Thus, these findings suggest that extinction learning of inhibitory avoidance engages a hippocampus-dependent consolidation process. Regarding contextual fear extinction, inconsistent results were reported. Fischer et al. (2004) showed that intra-hippocampus anisomycin injection immediately after extinction training improves contextual fear extinction without affecting auditory fear extinction. Mamiya et al. (2009) reported that this treatment does not alter the contextual freezing response extinction, whereas <u>De Carvalho Myskiw et al</u>. (2014) demonstrated that this inhibitor impairs extinction consolidation of the conditioned response.

The hippocampal protein synthesis that occurs during extinction of inhibitory avoidance seems to depend on gene expression triggered by the extinction process. It is blocked by pre-extinction training inhibition of hippocampal transcription whereas, as reported above, this inhibition does not affect the extinction when induced 1 or 3h after extinction training (Vianna et al., 2003). These findings contribute to demonstrate that extinction is indeed a form of associative learning and that it relies upon a single peak of transcription at the time of its acquisition. A recent report has also shown non-ribosomal protein synthesis involvement in the consolidation of contextual fear extinction. Non-ribosomal protein synthesis inhibitor rapamycin blocks fear extinction when given in the hippocampus after an extinction training session. This effect is not blocked by the co-administration of proteasome inhibitor  $\beta$ -lac which by itself is ineffective on extinction consolidation (De Carvalho Myskiw et al., 2014). However, Lee et al. (2008) reported that infusions of  $\beta$ -lac into the hippocampal CA1 region immediately after each extinction session suppress the extinction of this fear memory. They support the idea that extinction also involves some unlearning (or forgetting) process of the pre-existing contextshock association.

### 3.3. Cortex and other neural sites

In addition to the amygdala and hippocampus, the mPFC plays 1539 a crucial role in fear extinction (Tables 6 and 9). Lesion and inac-1540 tivation studies of mPFC reported contradictory results regarding 1541 its role in fear extinction (Akirav et al., 2006; Garcia et al., 2006; 1542 Laurent and Westbrook, 2008; Quirk et al., 2000; Sierra-Mercado 1543 et al., 2006). Such conflicting findings may be due to the fact that 1544 these studies have not distinguished between mPFC subregions. 1545 More recently, it was shown that the infralimbic cortex (IL), but not 1546 the prelimbic cortex (PL), is the subregion of the mPFC involved in 1547 fear extinction. Pre-extinction intra-IL, but not intra-PL, infusion of 1548 muscimol impaired extinction of auditory (Sierra-Mercado et al., 1549 2011) and contextual fear (Laurent and Westbrook, 2009). The IL 1550 appears to play different roles in two conditioned fear responses. It 1551 is implicated in extinction acquisition of freezing to an auditory CS, 1552 but not in its consolidation (Sierra-Mercado et al., 2011), whereas 1553 in contextual freezing extinction it is critical for consolidation and 1554 retrieval of this inhibitory learning (Laurent and Westbrook, 2009). 1555 Experiments performed on mPFC slices including IL from previ-1556 ously extinguished animals revealed that extinction training is 1557 associated with an increase of IL neurons excitability (Santini et al., 1558 2008). This effect is modulated by M-type potassium channels that 1559 contribute to the after-hyperpolarization that occurs after single 1560 action potential (Santini and Porter, 2010). In fact, the blockade of 1561 M-type K<sup>+</sup> channels in the IL before, but not after, extinction train-1562 ing facilitates auditory fear extinction, whereas activation of these 1563 channels before extinction training inhibits fear extinction (Santini 1564 and Porter, 2010). 1565

The role of other neural sites, both cortical and subcortical, in fear extinction was studied (Tables 6 and 9). The entorhinal cortex seems to be involved in this process. In our laboratory, it was shown that post-extinction training TTX blockade of ENT activity induces an impaired extinction retention of conditioned

### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

contextual freezing response, supporting the idea that ENT constitutes a critical component of neuronal network underlying fear extinction (Baldi and Bucherelli, 2014). Among the subcortical sites, we investigated the role of NBM and Substantia Nigra (SN) in contextual fear extinction. TTX inactivation of these neural sites immediately after extinction training does not affect subsequent extinction retention of conditioned freezing response (Baldi and Bucherelli, 2010). Thus, neither the NBM nor the SN are involved in extinction consolidation of fear memory. Little information is available regarding the role of sensory afferents to the forebrain, such as the thalamic nuclei, in the fear extinction. It was shown that the dorsal part of the midline thalamus containing mediodorsal, paraventricular and paratenial nuclei, is not necessary for auditory 1583 O3 fear extinction because its muscimol-induced inactivation before extinction training does not affect acquisition of extinction nor retention (Padilla-Coreano et al., 2012).

Expression of conditioned freezing is controlled by the midbrain periaqueductal gray (PAG) (LeDoux, 2000). This neural site was also implicated in extinction of freezing response to an auditory CS (McNally et al., 2004, 2005).

### 3.3.1. Neurotransmission systems

As in the amygdala, glutamatergic synaptic transmission within the mPFC contributes to fear extinction. Injection of intra-mPFC NMDA receptor antagonists performed pre- or post-extinction training impairs retention of fear conditioned responses extinction (Burgos-Robles et al., 2007; Fiorenza et al., 2012; Holmes et al., 2012), providing evidence that mPFC NMDA receptors are involved in fear extinction consolidation. Further support was obtained in experiments employing selective antagonist of NR2B-containing NMDA receptors. Pre-extinction administration of ifenprodil within the mPFC had no effect on fear conditioned responses (Laurent and Westbrook, 2008; Sotres-Bayon et al., 2009), whereas the same treatment applied immediately after extinction training impaired extinction retention (Laurent and Westbrook, 2008; Sotres-Bayon et al., 2009). To our knowledge there are few works investigating the effect of NMDA agonists directly injected into the mPFC. Fiorenza et al. (2012) reported that post-extinction administration of p-serine into the mPFC enhanced extinction retention of contextual fear and inhibitory avoidance responses. On the other hand, Chang and Maren (2011) found that pre-extinction training DCS infusion in this neural site does not facilitate auditory fear extinction, but enhances the subsequent reextinction fear. There are a few works analyzing the role of mPFC AMPA receptors. Zushida et al. (2007) observed extinction facilitation of contextual freezing response after pre-extinction injection of an AMPA "potentiator" in this cortical site. This effect is much more potent than that due to the intra-amygdala injection. Thus, the mPFC appears to be a major site in which AMPA "potentiator" acts enhancing fear extinction.

Recently it was proposed that in the mPFC NMDA receptor signaling is regulated by the voltage-gated calcium channels Cav2.1; this regulation is important for fear extinction. Using a contextual fear conditioning paradigm, Niimi et al. (2014) found that mice subjected to intracerebroventricular injections of Cav2.1 channels inhibitor after extinction training exhibit impaired extinction consolidation. This impairment is related to reduced Arc (CREB-dependent gene activity-regulated cytoskeleton-associated protein) expression in mPFC regions. Furthermore, transgenic mice carrying Cav2.1 gene mutation do not exhibit contextual freezing extinction when subjected to intra-mPFC injections of NMDA receptor antagonist (Niimi et al., 2014). Together these findings suggest that Cav2.1-mediated NMDA receptor signaling in the mPFC is involved in fear extinction consolidation.

Glutamatergic transmission within other cerebral sites appears to be involved in the fear extinction. The blockade of ENT NMDA receptors is followed by impaired extinction retention of the 1636 inhibitory avoidance response (Bevilaqua et al., 2006). Conversely, 1637 the same treatment administered in parietal and cingulate cortices 1638 or in the cerebellar nucleus interpositus does not affect extinction of 1630 inhibitory avoidance and fear-potentiated startle response, respec-1640 tively (Falls et al., 1992; Myskiw et al., 2010). These findings confirm 1641 that ENT, but neither parietal and cingulate cortices nor cerebel-1642 lum, are necessary for extinction to occur. Orsini and Maren (2009) 1643 reported that administration of NMDA or AMPA antagonists within 1644 the thalamic medial geniculate nucleus (MGN) before extinction 1645 training prevents extinction of conditioned fear, whereas neither 1646 protein synthesis inhibitor nor MAPK inhibitor affect this process. 1647 The authors suggested that the MGN is involved in auditory fear 1648 extinction as sensory information relay and it does not appear to 1649 be a locus of plasticity essential for formation of the extinction 1650 memory. 1651

Prefrontal GABAergic transmission is also involved in fear extinction. Local injections of GABAergic agonist muscimol impair fear memory extinction (Laurent and Westbrook, 2008, 2009; Sierra-Mercado et al., 2011, but see Akirav et al., 2006), and intramPFC infusion of GABAergic antagonist picrotoxin performed after extinction training facilitates extinction of auditory and contextual freezing response (Chang and Maren, 2011; Thompson et al., 2010). However, this effect is specific to the IL, as it is not observed if picrotoxin is administered into the PL (Thompson et al., 2010). These results further support a role of IL in fear extinction consolidation.

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Recent findings indicate that mPFC is a central site for noradrenergic modulation of extinction. Animals infused with the B-AR antagonist propranolol before, but not after, extinction training into the IL, exhibit impaired recall of extinction of auditory fear (Mueller et al., 2008). Similarly, the pre-extinction training administration of  $\alpha$ -ARs antagonist into the mPFC impairs acquisition of contextual conditioned fear (Do Monte et al., 2010). However, Fiorenza et al. (2012) showed that intra-mPFC injections of a different  $\beta$ -AR antagonist (timolol) immediately after an extinction session improve extinction retention of contextual fear, and impair that of inhibitory avoidance. In other words, β-AR blockade within mPFC has opposite effects on extinction of the two tasks. Moreover, norepinephrine administration in the same neural site and at the same time point induces impairment of contextual fear extinction, but has no effect on inhibitory avoidance extinction. Thus, modulation of fear extinction by the noradrenergic system into mPFC is complex.

Also, the mPFC expresses many dopaminergic receptors that 1679 may be involved in fear extinction modulation (Abraham et al., 1680 2014). Some studies have shown an increase of dopamine in this 1681 cortical site following fear extinction (Hugues et al., 2007) and 1682 decreased extinction retention after mPFC dopamine depletion 1683 (Espejo, 2003). Hikind and Maroun (2008) reported that D1 recep-1684 tors in the IL are involved in auditory fear extinction consolidation 1685 because pre- and post-extinction training injections of D1 antago-1686 nist result in an impairment of fear extinction. However, the admin-1687 istration of the same D1 antagonist in mPFC after extinction training 1688 of a contextual fear task has no effect on subsequent retention 1689 (Fiorenza et al., 2012). On the contrary, in an inhibitory avoidance 1690 task D1 antagonism impairs extinction consolidation (Fiorenza 1691 et al., 2012). Finally, D1 agonist injections after extinction session in 1692 the mPFC do not affect extinction retention of either contextual fear 1693 or inhibitory avoidance (Fiorenza et al., 2012). The D2 receptor has 1694 also been implicated in the modulation of fear extinction. Recently, 1695 it was shown that intra-IL administration of a selective D2 antago-1696 nist before extinction training does not affect extinction acquisition 1693 of auditory conditioned fear, but impairs extinction retention on 1698 the subsequent day, indicating the involvement of IL D2 receptor in extinction consolidation (Mueller et al., 2010). Moreover, this treat-1700 ment attenuates extinction-evoked firing in IL neurons (Mueller 170

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### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

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et al., 2010). Finally, D4 receptor blockade into the IL performed before auditory fear extinction induces normal extinction acquisition, but impaired extinction retention (Pfeiffer and Fendt, 2006), further suggesting that the dopaminergic activity in this cortical site is crucial for consolidation of fear extinction.

Also dopaminergic activity in the nucleus accumbens is relevant for fear extinction. Intra-accumbens pre-extinction administration of D2 receptors antagonist impairs both extinction acquisition and retention of conditioned freezing response to an auditory CS supporting the hypothesis that "accumbal dopaminergic activity regulates the development and retention of fear inhibition" (Holtzman-Assif et al., 2010).

A critical role in modulating fear extinction consolidation seems to be played by the cholinergic activity in the mPFC. Although, to our knowledge, only one study investigated this neurotransmitter system, it demonstrated that intra-IL blockade of muscarinic receptors before extinction training produces an impairment of auditory fear extinction. Scopolamine-treated animals exhibit normal extinction acquisition, but poor extinction memory retention (Santini et al., 2012). The authors speculated that in the IL a molecular mechanism underlying fear extinction might involve interactions between muscarinic cholinergic receptors and M-type K<sup>+</sup> channels (Santini et al., 2012).

Several studies reported findings supporting the role of CB1 receptors in fear extinction. Pre-extinction infusion of a CB1 antagonist within the IL blocks (Kuhnert et al., 2013; Lin et al., 2009), whereas IL infusion of CB1 agonist has no effect (Kuhnert et al., 2013) or enhances (Lin et al., 2009), extinction of fear-potentiated startle. The IL cannabinoid receptors appear also involved in the extinction of contextual conditioned fear (Do Monte et al., 2013a). The administration of cannabidiol (a non psychotropic phytocannabinoid) within this cortical site performed before each of three extinction sessions facilitates fear extinction. This facilitating effect is probably mediated by activation of IL CB1 receptors because systemic injection of a CB1 antagonist blocks this effect (Do Monte et al., 2013a).

Finally, as shown in the BLA, in the mPFC histaminergic system appears involved in fear memory extinction modulation. Experiments using contextual fear conditioning and the inhibitory avoidance paradigms revealed an extinction deficit in rats treated with the H2 receptors antagonist into the IL (Fiorenza et al., 2012). Because this effect was observed when the injections were performed immediately after extinction training, IL histaminergic receptors appear to modulate extinction consolidation. Conversely, intra-IL post-extinction infusion of histamine N-methyltransferase inhibitor had different effects on extinction of the two fear tasks: improved contextual fear extinction but had no effect on inhibitory avoidance (Fiorenza et al., 2012). Thus, the enhanced levels of histamine in the IL may have different effects on fear extinction consolidation depending on the fear memory task.

McNally and coworkers reported that pre-extinction training infusion into ventro-lateral PAG (vlPAG) of opioid receptors antagonist naloxone impairs auditory freezing response extinction; vlPAG opioid receptors play a specific role in the acquisition but not expression of extinction because the treatment fails to reinstate freezing to an already extinguished CS (McNally et al., 2004). Moreover, this effect is not observed following naloxone injections into dorsal PAG. These authors subsequently showed that vIPAG opioid receptors involved in fear extinction are  $\mu$  opioid receptors. In fact, intra-vlPAG administration of  $\mu$ -, but not  $\delta$ - or  $\kappa$ -opioid receptor antagonist retards auditory fear extinction (McNally et al., 2005). In addition, extinction is also impaired by intra-vlPAG injection of a cAMP analog suggesting that opioid antagonism effect in modulating fear extinction is mediated by cAMP inhibition that would occur with  $\mu$ -opioid receptor activation (McNally et al., 2005). Instead, administration of PKA activator or MAPK inhibitor within this neural site does not affect fear extinction (McNally et al., 2005).1768Finally, the authors found that intra-vlPAG pre-extinction training<br/>infusions of inhibitor of endogenous opioid catabolizing enzymes1769facilitate conditioned freezing response extinction to auditory CS<br/>(McNally, 2005) confirming a critical role for vlPAG endogenous<br/>opioids in fear extinction.1770

### 3.3.2. Protein kinases

MAPK, PKA and PI-3K within mPFC are critical for fear extinction, 1775 whereas CaMKII is not. It was reported that post-extinction inhibi-1776 tion of MAPK within this cortical site impairs extinction of auditory 1777 conditioned fear responses (Hugues et al., 2004). Moreover, phos-1778 phorylated MAPK is upregulated (Cannich et al., 2004; Kwapis et al., 1779 2014) and associated with enhanced levels of calcineurin into the IL 1780 after extinction training (Cannich et al., 2004). Mueller et al. (2008) 1781 found that intra-IL pre-extinction training injection of PKA antag-1782 onist, but not of CaMKII inhibitor, impairs subsequent extinction 1783 retention of auditory conditioned fear. Thus, PKA activity, but not 1784 CaMKII activity, is necessary for extinction consolidation within 1785 the IL. Finally, PI-3K inhibition within IL performed after extinction 1786 training hinders contextual fear extinction consolidation (Kritman 1787 and Maroun, 2013). 1788

Opposite results were obtained in ENT. In fact, intra-ENT infusion of CaMKII inhibitor impairs extinction of the inhibitory avoidance response, whereas the MAPK inhibitor administration has no effect (Bevilaqua et al., 2006). Together these findings confirm a crucial role of IL-mPFC and ENT in extinction consolidation of conditioned fear suggesting that different protein kinases are required in different brain structures.

### 3.3.3. Gene expression and protein synthesis

Several transcription factors into the mPFC are activated dur-1797 ing fear extinction. For example, Mamiya et al. (2009) reported 1798 an increased activation of CREB and CREB-dependent gene Arc 1799 within this neural site following extinction training of contex-1800 tual fear paradigm. Herry and Mons (2004) showed that auditory 1801 fear extinction is accompanied by an increase in c-fos and zif268 1802 expression into the mPFC; furthermore, resistance to re-extinction 1803 learning is associated with an impaired expression of these imme-1804 diate early genes in the same cortical region. These results were 1805 confirmed by other findings showing that rats selectively bred 1806 for high anxiety exhibit impaired extinction of auditory condi-1807 tioned fear response and low levels of c-fos expression within 1808 the IL (Muigg et al., 2008). Similarly, mice with specific extinction 1809 impairment also show decreased expression of intra-IL immediate 1810 early genes (Hefner et al., 2008). The transcription factor BDNF was 1811 also implicated in fear memory extinction. Using an auditory fear 1812 conditioning task Peters et al. (2010) reported that intra-IL pre-1813 extinction training infusion of BDNF facilitates extinction memory. 1814 Further confirmation of BDNF role in the IL in fear extinction was 1815 obtained by experiments showing that epigenetic modulation of 1816 BDNF genes in the IL is associated with auditory fear extinction 1817 (Bredy et al., 2007). The same authors recently reported that the 1818 activity of p300/CBP-associated factor within the IL is necessary for 1819 auditory fear extinction (Wei et al., 2012). 1820

Long-term memory for fear extinction requires new protein syn-1821 thesis in the ventral mPFC; rats subjected to pre-extinction training 1822 infusion of anisomycin within the IL exhibit normal extinction 1823 acquisition of conditioned freezing to an acoustic CS, but are unable 1824 to recall extinction the following day (Mueller et al., 2008; Santini 1825 et al., 2004). Similar results were obtained using a contextual fear 1826 paradigm and immediately post-extinction training administration 1827 (Mamiya et al., 2009). Moreover, Mueller et al. (2008) reported 1828 that in the IL the fear extinction consolidation-related protein 1829 synthesis dependents on new mRNA synthesis, because intra-IL 1830

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E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

pre-extinction training injection of the transcription inhibitor actinomycin impairs extinction retention.

Also protein synthesis in the ENT is important for fear extinction. The administration of anisomycin within ENT after the first extinction session of an inhibitory avoidance paradigm is followed by impaired extinction retention of these fear responses (Bevilaqua et al., 2006).

### 4. Discussion

The present review is centered on fear memory necessary for the organization of defensive behaviors and the survival of an organ-1840 ism. The role of several cerebral structures involved in fear memory reconsolidation and extinction was analyzed. In recent years there 1842 has been a growing interest in these two phases of fear memory processing, both induced by memory retrieval. During non-reinforced retrieval, a consolidated memory re-enters a vulnerable state during which it is again sensitive to disruption and, to persist, must undergo a new stabilization process (reconsolidation). The function of reconsolidation is a matter of debate. Two hypotheses have been proposed: (i) reconsolidation allows memory updating with new information, (ii) through reconsolidation the initial memory becomes stronger and longer lasting (Alberini and LeDoux, 2013; Tronson and Taylor, 2007). As the term reconsolidation is derived from consolidation, the mechanisms that underlie reconsolidation would be identical to those that mediate consolidation (Alberini, 2005; Dudai and Eisenberg, 2004). The two mnemonic processes seem to share some similar molecular mechanisms and pathways, such as protein synthesis, activation of MAPK pathway and the transcription factor CREB (Debiec et al., 2002; Dovere et al., 2007: Mamiya et al., 2009; Nader et al., 2000; Tronson and Taylor, 2007). 1859 Nevertheless, it was shown that reconsolidation is not an exact 1860 recapitulation of consolidation; the two processes show different time courses (reconsolidation is completed faster than consolidation) and differences at the neural circuits and molecular levels (Alberini and LeDoux, 2013; Mactutus et al., 1979; Tronson and Taylor, 2007). This is not surprising considering the procedural differences involving the two processes. Indeed, consolidation is induced only presenting the CS and US contiguously, whereas reconsolidation is induced presenting either the CS or the US alone.

Reconsolidation would be a behavioral phenomenon opposing extinction, the classical retrieval-induced process caused by changes in the associative relationships that generated the original response. Extinction is not oblivion because the original response recovers spontaneously over time, presenting the CS in a new context (renewal) and upon unpredictable US presentations (reinstatement) (Myers and Davis, 2007; Quirk and Mueller, 2008). These behavioral properties indicate that during extinction a new inhibitory memory trace is formed that competes with the original fear memory (Myers and Davis, 2007; Myskiw et al., 2014; Pape and Pare, 2010; Quirk and Mueller, 2008). Reconsolidation and extinction processes are operationally similar. Both phases are induced by non-reinforced presentation of the CS (Nader and Hardt, 2009; Quirk and Mueller, 2008), but they have opposing actions on the fate of the retrieved memory. Reconsolidation stabilizes or strengthens the memory trace, whereas extinction induces new opposite learning. In other words, reconsolidation and extinction are competing processes. The competition between them seems to depend partly on the length and/or number of memory reactivation sessions. A brief re-exposure, like that caused by a short retrieval session, would induce reconsolidation, whereas longer or repeated reminder trials would result in extinction (Debiec et al., 2002; Eisenberg et al., 2003; Pedreira and Maldonado, 2003). The growing interest about these memory phases is witnessed by the exponential increase in publications related to the two phenomena

(Besnard et al., 2012; Delamater and Westbrook, 2014). This is 180/ due in part to the fact that understanding the mechanisms of fear 1805 memory reconsolidation and extinction may offer new therapeutic 1806 interventions for the treatment of human fear and anxiety disor-1807 ders, such as phobias and post-traumatic stress-disorder (PTSD) 1808 characterized by dysregulated fear responses (Alberini, 2005; 1800 Auber et al., 2013; Davis et al., 2006; Hartley and Phelps, 2010; 1900 Monfils et al., 2009; Nader, 2003; Parsons and Ressler, 2013; Quirk 1001 et al., 2010; Rao-Ruiz et al., 2011; Rossato et al., 2010; Schiller 1902 et al., 2010; VanElzakker et al., 2014). Thus, the identification of 1903 both neural structures and molecular mechanisms underlying the 1904 two memory phases appears to be crucial. 1905

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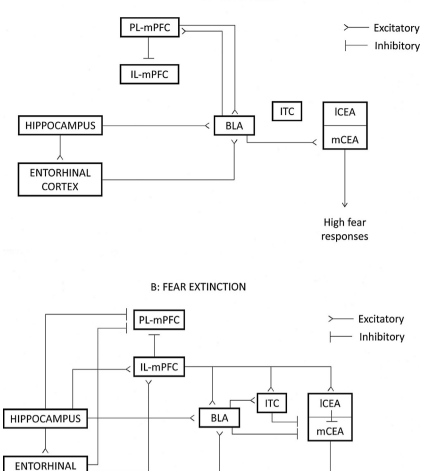
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### 4.1. Neural circuit underlying fear memory reconsolidation and/or extinction

Reconsolidation and extinction show different anatomical and 1908 biochemical signatures (Mamiya et al., 2009; Merlo and Romano, 1909 2008; Suzuki et al., 2004). Although the experimental results are 1910 not always consistent, the amygdala and hippocampus appear to 1911 be the neural sites playing a key role in both reconsolidation and 1912 extinction of fear memory. Yet, whereas the amygdala is involved 1913 in these memorization phases whatever fear memory task is con-1914 sidered (cued and contextual fear conditioning, fear potentiated 1915 startle or inhibitory avoidance), the hippocampus is involved when 1916 contextual components are implicated. In addition to these brain 1017 structures, evidence points to a crucial role of the mPFC in these 1018 mnemonic processes. However, fear memory reconsolidation and 1010 extinction seem to involve different subregions of this cortical site. 1920 Indeed, the PL region appears to be implicated in reconsolidation 1921 of conditioned fear responses, whereas the IL region appears to be 1922 the candidate to suppress fear responses via extinction learning. 1923 Finally, recent results have shown that the ENT as well might be part 1924 of a circuit underlying fear memory reconsolidation and extinction. 1925 These neural sites are closely interconnected. Anatomical studies 1926 revealed that there are reciprocal connections between the amyg-1927 dala, the hippocampus and the ENT. The hippocampal CA1 field 1928 and the ENT are among the prominent sources of amygdalar affer-1929 ents and project mainly to the BLA (Canteras and Swanson, 1992; 1930 Ottersen, 1982; Pitkanen et al., 2000; Wyss, 1981). The BLA in turn 1931 projects abundantly to the hippocampus (with dense synapses on 1932 the CA1 field) and ENT (Pikkarainen et al., 1999). ENT provides the 1933 major gateway for transmission of information between the hippocampus and cortex (Hyman et al., 1990; Maren and Fanselow, 1935 1997). Several lines of evidence suggested that the amygdala mod-1936 ifies the hippocampus and ENT responses and vice versa (Abe, 1937 2001; Maren and Fanselow, 1995; McGaugh, 2000; Packard and 1938 Cahill, 2001; Richter-Levin and Akirav, 2000). This seems to be 1939 also supported by experimental findings that theta synchrony of 1940 hippocampal CA1 and LA increases and decreases during fear mem-1941 ories reconsolidation and extinction, respectively (Narayanan et al., 1942 2007; Sangha et al., 2009). Moreover, it was reported that BLA and 1943 ENT neuronal activity oscillates in phase (Pare and Gaudreau, 1996; 1944 Paz and Pare, 2013) both structures interacting in the modula-1945 tion of fear memory consolidation (Roesler et al., 2002). Thus, a 1946 dynamic interaction may exist between the amygdala, hippocam-1947 pus and ENT underlying the dynamic nature of memory processes. 1948 The PL and IL are strongly interconnected with each other (Hoover 1949 and Vertes, 2007) and with the amygdala, hippocampus and ENT. 1950 The PL and IL receive massive afferents from hippocampal CA1 1951 field and ENT, and in turn send projections to these same neural 1952 sites (Hoover and Vertes, 2007; Vertes et al., 2007). It was reported 1953 that fear extinction is related to LTP-like synaptic changes in DH-1954 mPFC projection; low frequency stimulation of the DH attenuates this synaptic plasticity and impairs extinction retention, whereas high frequency stimulation of the DH has opposite effects (Farinelli

E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

### A: FEAR RECONSOLIDATION



**Fig. 1.** Proposed neural circuits underlying fear memory reconsolidation and extinction. (A) During fear memory reconsolidation, hippocampal and entorhinal inputs enhance the basolateral amygdala (BLA) activity that excites pyramidal neurons of the prelimbic region within the medial prefrontal cortex (PL-mPFC). In turn, the PL-mPFC input synapses on infralimbic region of the mPFC (IL-mPFC) inhibiting it and on the BLA (probably on "fear neurons") which influences central amygdala activity (CEA), the amygdaloid output nucleus. The result is to rise conditioned fear response. (B) During fear memory extinction, hippocampal and entorhinal inputs inhibit the PL-mPFC, whereas excite the BLA and IL-mPFC. In turn, the IL-mPFC contributes to inhibit PL-mPFC and may act by exciting (i) the BLA "extinction neurons" which inhibit the medial division of CEA (mCEA), directly or indirectly (through intercalated mass cells, ITCs), (ii) GABA-ergic ITCs that inhibit mCEA and (iii) inhibitory interneurons within the lateral division of CEA (ICEA) that, in turn, inhibits mCEA. The result is to lower conditioned fear response.

et al., 2006). Moreover, the IL is the primary site of action for hip-1958 pocampal BDNF and increasing BDNF in this pathway prevents fear 1959 extinction impairment (Peters et al., 2010). Thus, hippocampus-IL 1960 projection appears to be a key projection for fear memory extinc-1961 tion. Sotres-Bayon et al. (2012) suggested that following extinction 1962 an increased inhibition of PL activity takes place. This might be due 1963 partly to the hippocampal inputs that excite local PL interneurons 1964 triggering feed-forward inhibition of PL neurons (Sotres-Bayon 1965 et al., 2012) and in part to the inhibitory actions of IL on PL (Ji and 1966 Neugebauer, 2012). These two regions of the mPFC project differ-1967 **04** ently to the amygdala. Whereas PL fibers selectively target the BLA 1968 and CEA, IL fibers are distributed mainly to medial and basome-1969 dial nuclei of the amygdala, intercalated cell masses and lateral 1970 division of CEA (ICEA) (Vertes, 2004). It is likely that differential 1971 activation of the two regions of the mPFC and consequently of their 1972 differential connectivity with the amygdalar nuclei orchestrate 1973 conditioned fear responses during reconsolidation and extinction 1974 1975 processes. Supposedly, during fear reconsolidation enhanced BLA activity driven by hippocampal and entorhinal inputs controls 1976

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PL activity triggering PL pyramidal neurons. PL input synapse on 1977 "fear neurons" within the amygdaloid nuclei, which fire selectively 1978 during and after fear conditioning (Herry et al., 2008) (Fig. 1A). 1979 These neurons, in turn, may influence CEA activity thus modu-1980 lating the expression of conditioned fear responses by means of 1981 projections to midbrain and hypothalamic sites or the ventrolat-1982 eral PAG (freezing) (LeDoux, 2000). On the contrary, during fear 1983 extinction, PL activity could be inhibited (due to the stimulation 1984 of the PL interneurons by hippocampal and IL projections) (Ji and 1985 Neugebauer, 2012; Sotres-Bayon et al., 2012;) whereas IL activity 1986 is stimulated (Ji and Neugebauer, 2012; Knapska et al., 2012; Milad 1987 and Quirk, 2002). IL inputs may synapse on "extinction neurons" 1988 within amygdalar nuclei, which fire selectively to an extinguished 1989 CS (Herry et al., 2008). Extinction neurons may then inhibit the out-1990 put of CEA (Fig. 1B). Alternatively, or additionally, IL excitatory out-1991 put may activate the ITC neurons that in turn inhibit CEA providing 1992 a mechanism of extinction (Amano et al., 2010, 2012; Ehrlich et al., 1993 2009; Likhtik et al., 2008; Pape and Pare, 2010; Pare and Duvarci, 1994 2012; Quirk and Mueller, 2008). Moreover, the ITC neurons might 1995

Low fear responses

#### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

integrate additional inputs from the BLA to set the level of inhibition of CEA neurons (Amano et al., 2011) (Fig. 1B). Finally, IL-ICEA projections might activate inhibitory interneurons within ICEA which in turn inhibit output neurons of mCEA (Fig. 1B). Thus, although fear reconsolidation and extinction involve the same neural structures, they may take place in distinct neuronal circuits involving different subregions, connections and neuronal populations. This seems to be supported by electrophysiological and immunohistochemical experiments that have identified distinct amygdaloid ("fear neurons" and "extinction neurons") and hippocampal ("cFos<sup>+</sup> cells" and "pERK<sup>+</sup> cells") neurons activated during conditioning and extinction of fear (Herry et al., 2008; Tronson et al., 2009). Moreover, the two subpopulations of amygdalar neurons have preferential connections either to the PL or the IL; whereas the amygdaloid neurons whose activity is correlated with fear memory are innervated by the PL, those whose activity is correlated with fear extinction receive inputs mainly from the IL (Ji and Neugebauer, 2012; Knapska et al., 2012). Therefore, amygdaloid fear neurons and hippocampal cFos<sup>+</sup> cells might be connected with ENT and PL projections that are activated during reconsolidation of conditioned fear responses, whereas amygdaloid extinction neurons and hippocampal pERK<sup>+</sup> cells might be connected with ENT and IL projections activated during extinction of these fear responses. Thus, fear memories reconsolidation and extinction are two competing mnemonic phases which require the activation of different neuronal circuits.

4.2. Temporal and biochemical signatures in the neural sites involved in reconsolidation and extinction

The diverse anatomical requirements presumably are related 2023 to the distinct temporal and biochemical signatures. Fear memo-2024 ries reconsolidation and extinction have different temporal profiles 2025 (Tronson et al., 2012). After retrieval, there is a brief time window 2026 for reconsolidation, whereas extinction only takes place after pro-2027 longed re-exposure to the CS in absence of the US (Suzuki et al., 2028 2004). At the molecular level, the activity of several molecules is 2029 required for both processes, but others are oppositely regulated 2030 during the two phases (De la Fuente et al., 2011; Merlo et al., 2031 2005; Merlo and Romano, 2008). For example, both fear mem-2032 ory reconsolidation and extinction are protein synthesis dependent 2033 processes, as shown by their disruption when a protein synthesis 2034 inhibitor is administered after memory reactivation or extinction 2035 training (see Tables 1–3 and 7–9), but the protein synthesis may 2036 require different upstream receptors, signaling and transcription 2037 2038 factors. For example, increased levels of phosphorylated GluR1 subunit-containing AMPA type glutamate receptor were found 2039 in the lateral amygdala after fear memory reactivation, whereas 2040 its dephosphorylation was observed after fear memory extinction 2041 (Monfils et al., 2009). It was also shown that activation of the endo-2042 cannabinoid system reduces the reconsolidation of fear memories, 2043 whereas its hypo-activation promotes their reconsolidation lead-2044 ing to enduring fear responses (De Oliveira Alvares et al., 2008; 2045 Lin et al., 2006). On the contrary, intact CB1 receptor signaling 2046 appears to be essential for proper extinction of aversive mem-2047 2048 05 ories (Abush and Akirav, 2010; De Oliveira Alvares et al., 2008; Ganon-Elazar and Akirav, 2009; Kunhert et al., 2013; Lin et al., 2049 2009). Therefore, it may be postulated that the endocannabinoid 2050 system determines the balance between the processes of main-2051 taining or strengthening the original memory (reconsolidation) 2052 and the establishment of a new memory (extinction) (De Oliveira 2053 Alvares et al., 2008). Both processes require NMDA receptors acti-2054 vation whereas fear extinction, but not reconsolidation, involves 2055 L-VGCCs (Davis and Bauer, 2012; De Carvalho Myskiw et al., 2014; 2056 Suzuki et al., 2008). Increased intracellular calcium results in the 2057 2058 protein kinases activation, such as MAPK, that translocate into the 2059 nucleus where they activate (phosphorylate) several transcription factors to promote gene transcription and new protein synthesis. 2060 The two isoforms of MAPK, ERK1 and ERK2, seem to be involved 2061 in a different manner in the two mnemonic phases (Cestari et al., 2062 2014). Indeed, whereas fear reconsolidation primarily involves 2063 ERK2 (Cestari et al., 2006), an increased intranuclear pERK1 has 2064 been reported during fear extinction (Fischer et al., 2007). Also 2065 PI-3K and its downstream target AKT seem to be recruited in 2066 different way in fear memories reconsolidation and extinction: 2067 they are reactivated and dephosphorylated, respectively (Lin et al., 2068 2003a). Furthermore, the outcome of retrieval in terms of recon-2069 solidation/extinction may depend on the balance between protein 2070 kinases and phosphatases (such as calcineurin) activity. As it has 2071 been proposed by Lin et al. (2003a), the stimulation of MAPK 2072 may activate several transcriptional factors to reactivate origi-2073 nal memory on one hand and promote calcineurin synthesis on 2074 the other hand. Calcineurin, in turn, may exerts a negative feed-2075 back effect to down-regulate kinases. Therefore, when protein 2076 kinases activity dominates the reconsolidation process is trig-2077 gered, when calcineurin activity dominates the extinction process 2078 is triggered. Finally, fear memory reconsolidation and extinction 2079 may involve either different transcription factors or the same transcription factors but in different manner. For example, both processes involve NFkB, yet activity of the transcription factor NFAT 2082 is engaged by extinction but not reconsolidation (De la Fuente et al., 2083 2011).

As previously mentioned, extinction is a form of new learning 2085 and as such it consists of an acquisition and a consolidation phase 2086 (Myers and Davis, 2007; Pape and Pare, 2010; Quirk and Mueller, 2087 2008). The two phases are usually studied by means of treatments 2088 applied pre- or post-extinction training, respectively. Although it is 2089 not completely understood how the several brain sites contribute 2090 to each phase of extinction process, the results tend to support the 2091 involvement of the BLA in both phases and the mPFC, hippocam-2092 pus and ENT only in the consolidation phase (Baldi and Bucherelli, 2093 2014; Bevilagua et al., 2006; Quirk and Mueller, 2008). For example, 2094 fear extinction acquisition activates amygdalar NR2B-containing 2095 NMDA receptors that induce calcium influx. During extinction con-2096 solidation in the same structure L-VGCCs are activated allowing 2097 further increase of intracellular calcium concentration, whereas in 2098 mPFC the NR2B are activated, perhaps following stimulation by the 2099 amvgdalar inputs. 2100

### 4.3. *Reconsolidation and extinction in human anxiety disorders*

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As mentioned above, understanding the biological mechanisms 2102 of fear memory reconsolidation and extinction may have clinical 2103 relevance in treating human anxiety disorders such as PTSD. PTSD 2104 patients show strong traumatic memories that are continuously 2105 retrieved in an intrusive manner, causing re-experiencing of the 2106 traumatic event and increased arousal and stress response. The 2107 persistence of PTSD can be explained in terms of trauma-induced 2108 strengthening of the memory trace or failure to extinguish con-2109 ditioned fear memory (Alberini and LeDoux, 2013; VanElzakker 2110 et al., 2014). Thus, the pharmacological interferences effective in 2111 disrupting fear memory reconsolidation or enhancing extinction 2112 could potentially be useful for reducing expression of fear mem-2113 ory (Fitzgerald et al., 2014; Quirk and Mueller, 2008). Based on 2114 results obtained in rodents, translational studies in humans are 2115 beginning to be carried out. For example, the  $\beta$ -adrenergic antago-2116 nist propranolol and mTOR blocker rapamycin could be promising 2117 treatments for targeting the fear memory reconsolidation. The 2118 oral administration of propranolol before fear memory reactivation 2119 in healthy human subjects reduced significantly fear-potentiated 2120 startle response during testing 24 h later and prevented the return 2121 of fear (Kindt et al., 2009). Moreover, the same pharmacological 2122 treatment in patients suffering from PTSD reduced physiological 2123

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parameters of fear when the subjects again described their traumatic experience a week later (Brunet et al., 2008). Recently, rapamycin combined with reactivation of a traumatic memory was used in a pilot study in male veterans. The results showed that veterans treated with rapamycin (sirolimus) reported significantly fewer and less intense PTSD symptoms 1 month later, although the effects did not persist at 3 months (Suris et al., 2013).

In the treatment of PTSD exposure-based therapy is frequently 2131 used. It is conceptually based upon fear extinction. DCS is the best 2132 studied extinction enhancer and has been used as an adjunct to 2133 psychotherapy in humans (Davis, 2011; Hofmann et al., 2013b). 2134 In clinical studies, DCS administered before the exposure ses-2135 sions improves responses to therapies for acrophobia (fear of 2136 heights, Ressler et al., 2004), social anxiety disorder (Hofmann 2137 et al., 2013a) and panic disorder (Otto et al., 2010); however, it 2138 seems to be less effective in therapeutic treatment of PTSD (De 2139 Kleine et al., 2012; Litz et al., 2012). In rodents, other drugs were 2140 shown to facilitate extinction and might be useful in humans. These 2141 include glucocorticoids and cannabinoid agonists. PTSD patients 2142 have reduced circulating levels of cortisol (Yehuda, 2001) and it 2143 has been shown that glucocorticoids affect symptoms severity. In 2144 2145 fact, hydrocortisone administration enhances exposure therapy in PTSD (Suris et al., 2010; Yehuda and LeDoux, 2007) and low-dose 2146 cortisol improves treatment of PTSD symptoms (Aerni et al., 2004). 2147 Cannabinoid agonists also facilitated fear extinction memory in 2148 healthy humans (Rabinak et al., 2013) and this effect appeared 2149 to be due to the modulation of prefrontal-hippocampal circuits 2150 (Rabinak et al., 2014). However, these agents are not yet utilized 2151 in the treatment of anxiety disorders. 2152

Reconsolidation and extinction might interact at both pharma-2153 cological and procedural levels. Their pharmacological interaction 2154 may constitute a limit for the use of the reactivation or expo-2155 sure therapy for the treatment of anxiety disorders in humans. 2156 Indeed, the use of exposure to cues to retrieve and extinguish fear 2157 memories could, under some circumstances, result in strengthen-2158 ing of fear memory. This is important when extinction-enhancing 2159 agents (such as DCS) or reconsolidation-impairing drugs (such 2160 propranolol) are used. DCS accelerates and strengthens fear extinc-2161 tion, but it also enhances fear memory reconsolidation (Lee et al., 2162 2006). Similarly, propranolol impairs fear memory reconsolida-2163 tion but also impairs fear extinction resulting in high fear (Cain 2164 et al., 2004). The result might be a potentially strengthening of 2165 maladaptive memories after retrieval. Because the duration of the 2166 re-exposure to the CS appears to be an important factor, the phar-2167 2168 macological agent used must be coordinated with the exposure duration for targeting the right memory phase. On the other hand, 2169 procedural interactions between reconsolidation and extinction 2170 might be an alternative to pharmacological intervention for the 2171 treatment of anxiety disorders. Recently, several studies demon-2172 strated that fear extinction performed during a reconsolidation 2173 window enhances the effects of extinction training preventing 2174 the re-expression of fear memory in rodents (Auber et al., 2013; 2175 Monfils et al., 2009; Pineyro et al., 2014; Quirk et al., 2010; Rao-Ruiz 2176 et al., 2011; Rossato et al., 2010). It was proposed that mak-2177 ing the original memory labile through reactivation, extinction 2178 learning overwrites the original memory (Monfils et al., 2009). 2179 Similar results have been reported in humans. Schiller et al. 2180 (2010) showed that post-retrieval extinction may interfere with 2181 the fear memory reconsolidation in humans and it selectively 2182 blocks the reconsolidation of the retrieved memory but does not 2183 affect non-retrieved memories. On the contrary, Soeter and Kindt 2184 (2012) failed to replicate these results using different fear memory 2185 responses. 2186

In conclusion, although the inconsistent findings indicate the need for further investigation, the improvement of these interventions could lead to new therapeutic treatments of 2189 pathological fear memories. 2190

Uncited references

Huang et al. (2013), Iwata et al. (1986), LeDoux et al. (1986), LeDoux et al. (1985) and LeDoux et al. (1983).

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### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

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### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

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#### E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

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E. Baldi, C. Bucherelli / Neuroscience and Biobehavioral Reviews xxx (2015) xxx-xxx

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