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Histaminergic neurotransmission as a gateway for the effects of the fat sensing molecule Oleoylethanolamide

Focus on cognition and stress-reactivity

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Abstract

The first description of histamine dates back to Sir Henry Dale and co-workers (Dale HH, 1910; Kwiatkowski, 1941) who detected its presence and biological activity. Dale and colleagues by studying the effect of ergot extracts, identified histamine as stimulant of smooth muscle in the gut and respiratory tract by induction of vasodilation. Simultaneously, at the beginning of XX century, the stimulation of acid secretion in the stomach caused by histamine was also recognized. During the first 20 years after these discoveries other important histaminergic actions were defined. Lewis and Grant (LEWIS T., 1924) headlines: Vascular reaction of the skin to injury. The liberation of a histamine-like substance in injured skin: the underlying cause of factitious urticarial and of wheals produced by burning, and observation upon the nervous control of skin reactions. With this discovery was defined for the first time what will be then called "triple response of Lewis". The rapid developments of these years soon ceased, however, and despite continuous efforts and considerable improvements in the available technical facilities the experimental data since presented to show that histamine participates in physiological processes appear to be inconclusive. The presence of histamine in the brain, predominantly in the grey matter, was first shown by Kwiatkowski (Kwiatkowski, 1941), and White (WHITE, 1959) but in 1960's with the introduction of fluorimetric assay that disclosed the anatomical identity of the catecholamine containing neurons and their projections, histamine was initially neglected because the assay failed to determine the location of histamine in the brain. This was because the reagents that were used to detect this diamine cross-reacted with spermidine—a uniformly distributed polyamine that occurs at high concentrations (Haas and Panula, 2003). But circumstantial evidence began to accumulate for a significant role for histamine, and lesion studies by the Schwartz group, in 1974, pointed to the approximate location of a brain source of histamine (Garbarg et al., 1974). Later, two different and parallel discoveries brought to a general acceptance of the existence of central histaminergic system. Panula and colleagues and Watanabe and co-workers in 1984 independently demonstrated by means of immunohistochemistry that the tuberomamillary nucleus (TMN) in the posterior hypothalamus, is the only source of histaminergic neurons and the origin of the widely distributed histaminergic projections just like the other amine systems (Panula et al., 1984; Takeda et al., 1984). It is now clear that the histaminergic TM system commands general states of metabolism like satiety and appetite, arousal, and of consciousness, including learning and memorizing both pleasurable and aversive events.

In my thesis, I will provide evidence that in the brain histaminergic system converge peripheral and central signals and orchestrates appropriate behavioural responses. In particular, I will present results that strongly suggest the involvement of
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In my thesis, I will provide evidence that in the brain histaminergic system converge peripheral and central signals and orchestrates appropriate behavioural responses. In particular, I will present results that strongly suggest the involvement of

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Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide histamine neurons in the pro-cognitive and antidepressant-like effects of oleoylethanolamide, a lipid that is normally secreted by the intestine after a fatty meal, and I will present data on some of the molecular mechanisms responsible for these effects. I will also illustrate the influence of dietary factors on stress reactivity and cognition and preliminary results on the role of brain histamine in such a model.

The results exposed in this thesis may have relevance in revealing new mechanisms in the gut brain axis when brain histamine plays a key role as it acts as a gatekeeper of peripheral and central signals to elaborate the appropriate behaviour.
Chapter 1
Histamine in the Nervous System

1.1 Anatomic Framework

In the central nervous system, the presence of histamine is attributed at the presence of the histamine-releasing neurons. The amount of histamine derived by non-neuronal pool (mast cells) is somewhat limited under normal conditions. Other possible sources of histamine in the brain may include microglia and microvascular endothelial cells (Katoh et al., 2001; Yamakami et al., 2000). Mast cells are relatively scarce in the brain, in comparison to other tissues, and their function is at present unclear. Furthermore, the amount of peripherally synthetized histamine not contribute to its central content due to the histamine inability to cross the blood brain barrier. Therefore, it can be assumed that central histaminergic function are due almost exclusively to histaminergic neurons (Brown and Ennis, 2001). As previously reported, the histamine-producing neurons are located in the small tuberomamillary nucleus (TMN). The name, tuberomamillary nucleus, derives from the anatomical term tuber cinerum, denoting an ashen swelling located rostral to the mammillary bodies and caudal to the optic chiasm, forming the floor of the third ventricle in the hypothalamus (Krüger and Nyland, 1995). The TM in rats has been subdivided by Ericson et al. (Ericson et al., 1987) into three subgroups: (I) the medial tuberomamillary subgroup (TMM), which consists of around 600 neurons located on either side of the mammillary recess; (II) the ventral tuberomamillary subgroup (TMV), which contains approximately 1500 neurons around the mamillary bodies; and (III) the diffuse part of the TM (TMdiff or E5), which is made up of about 100 HD-immunoreactive perikarya scattered within or between various hypothalamic nuclei (Inagaki et al., 1990). In the mouse brain the TMN is less compact and is characterized by smaller and fewer neuron than rat TMN (Parmentier et al., 2002). The human TMN consist of about 64.000 neurons anatomically identified as the ventral, medial area and the lateral area (Airaksinen et al., 1991a).

The histamine neurons in the TM send projections that innervate the entire brain, and parts of the spinal cord (Figure 1) (Panula et al., 1984; Watanabe et al., 1984). Two ascending pathways and one descending pathway have been identified (Panula et al., 1989). The highest density of histaminergic fibres are found in the hypothalamus, diagonal band, septum and olfactory tubercle. Moderate density of fibres are found in cerebral cortex, striatum and nucleus accumbens. Projections to the midbrain, brain stem, cerebellum and spinal cord tend to be of lower density. The hippocampal formation is most strongly innervated in the subiculum and dentate gyrus, with a low...

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Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide density of fibres present in CA3 and CA1 (Brown and Ennis, 2001). The afferent projections to TM neurons are widespread and come from many different areas. Prominent sources are the infralimbic prefrontal cortex, lateral septum and preoptic nucleus (Ericson et al., 1991). Most of the efferent histaminergic fibers are unmyelinated and except for those that project to the trigeminal nucleus (Inagaki et al., 1988), do not in general form synaptic specializations, rather, histamine is released from varicosities located periodically along the axon (Takagi et al., 1986). Thus, histamine release sites and histamine receptors are not directly associated to one another. Rather, histamine has been proposed to act like a local hormone on neurons, glial cells and blood vessels in a concerted manner (Wada et al., 1991). In addition to histamine, the TM neurons contain several other neurotransmitters and modulators like GABA that is presumably released in specific brain regions to modulate behavioural response (Williams et al., 2014; Yu et al., 2015b). The neuropeptides GALANIN, thyrotropin-releasing hormone, proenkephalin-derived peptides and substance P are also found in various populations of histamine producing TM neurons. Most TM neurons also express adenosine deaminase (ADA), the enzyme that catalyzes the conversion of adenosine to inosine (Haas and Panula, 2003).

1.2 Histaminergic Receptors

The basic homeostatic and higher functions, including cognition, arousal, circadian and feeding rhythms regulated by brain histamine are due to the action on 4 metabotropic receptors: H1R, H2R, H3R, H4R. All of histaminergic receptors are expressed at central level with different density in different brain regions (Passani and Blandina, 2011). All metabotropic histamine receptors (H1R-H4R) belong to the rhodopsin-like family of G protein coupled receptors (GPCR). Each receptor consists of seven large transmembrane-spanning elements with prototypic domains (Haas et al., 2008). Three of the four histamine receptors that have been identified (H1–H3) are prominently expressed in the brain in specific cellular compartments, whereas the fourth (H4) receptor is detected predominantly in bone marrow and leukocytes (Haas and Panula, 2003).

1.2.1 Histamine H1 Receptor

The human H1 receptor is encoded by a gene of 56kDa composed by 487–490 amino acids located on chromosome 3p25 (Jongejan et al., 2005). The signal transduction of H1R is mainly mediated by coupling to Gq/11 proteins (Gutowski et al., 1991; Leopoldt et al., 1997; Moniri et al., 2004; Selbach et al., 1997), but also signals via G/0 in some systems (Seifert et al., 1994; Wang and Kotlikoff, 2000), and the small G protein family, most likely through an indirect downstream effect (Mitchell and Mayeenuddin, 1998). The interaction of H1 receptor with Gq/11 protein and phospholipase C promotes inositol trisphosphate (IP3)-dependent Ca2+ release from intracellular Ca2+-stores, and also diacylglycerol formation. H1R also activates AMP-kinase, nuclear factor kappa B, nitric oxide synthases, and phospholipase A2 (PLA2), which induces arachidonic acid formation (Haas et al., 2008). H1R are found throughout the whole body and nervous system. H1 receptors are widely distributed in mammalian brain (Hill, 1990; Schwartz et al., 1991). High densities are found in brain
regions concerned with neuroendocrine, behavioural, and nutritional state control, including the periventricular, suprachiasmatic, and ventromedial nuclei of the hypothalamus, aminergic and cholinergic brainstem nuclei, thalamus, and cortex (Schwartz et al., 1991). The global loss of H1R in KO mice produces immunological, metabolic, and behavioural abnormalities (Haas et al., 2008; Hirai et al., 2004; Huang et al., 2006; Masaki and Yoshimatsu, 2006).

1.2. Histamine H2 Receptor

A second class of histamine receptors was identified by Black and colleagues based on the different pharmacological profile of the histamine receptor responsible for stimulating gastric acid secretion (Hill et al., 1997). The gene encoding the human H2R, which is a 40-kDa 359-amino acid peptide, is located on chromosome 5q35.5. H2R couple to Gs proteins to stimulate adenylyl cyclase and increase intracellular cAMP, which activates protein kinase A (PKA) and the transcription factor CREB, all of which are key regulators of neuronal physiology and plasticity. Through H2R activation and PKA-dependent phosphorylation, histamine blocks a Ca2+-activated potassium conductance responsible for the neuronal excitability (Haas et al., 2008). Independent of either cAMP or [Ca2+]i levels, H2R also inhibit PLA2 and release of arachidonic acid, which likely account for the opposing physiological responses elicited by H1R and H2R in many tissues (Traiffort et al., 1992). Like the histamine H1 receptor, the H2 receptor has a widespread expression in the brain and spinal cord, particularly high densities are found in the basal ganglia and in parts of the limbic system such as the hippocampal formation and amygdala. In contrast to H1 receptors, H2 receptors are present in low densities in septal areas, hypothalamic and thalamic nuclei. H1 and H2 receptors are colocalized in several areas of the brain including pyramidal and granule cells in the hippocampal formation and in the other aminergic cell where the receptors can act synergistically, e.g. in the stimulation of cAMP production (Brown and Ennis, 2001). Mice deficient in H2R function exhibit selective cognitive deficits along with an impairment in hippocampal LTP (Dai et al., 2007) and with abnormalities in nociception (Mobarakhe et al., 2006; Mobarakhe et al., 2005) and gastric and immune functions (Teuscher et al., 2004).

1.2.3 Histamine H3 Receptor

Histamine H3 receptor in the brain were detected in1983 by the group of J.C. Schwartz in Paris proved its neurotransmitter function as auto- as well as hetero-receptor at pre- and postsynaptic membranes and revealed its profound influence on different neurotransmitter balances (Panula et al., 2015). The gene (Hrh3), encoding human H3R, a 70-kDa 445-amino acid peptide, is located on chromosome 20q13.33. H3R negatively couple through pertussis toxin-sensitive Gi/o proteins to N- and P-type Ca2+ channels and to adenylyl cyclase. Through extensive cross-talks with other GPCRs, they can also engage Gq/11 signaling and activate PLA2, Akt/GSK3, and MAP kinase pathways, all of which play important roles in axonal and synaptic plasticity and a variety of brain disorders (Haas et al., 2008). The histamine H3 receptor is located on histaminergic neuron somata, dendrites and axon varicosities, as well as on
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the axon varicosities and somata of other neurons, providing negative feedback to
inhibit histamine synthesis and the release of histamine or other transmitters, including
glutamate (Brown and Reymann, 1996; Doreulee et al., 2001), acetylcholine
(Arrang et al., 2007; Passani and Blandina, 1998) and GABA (Jang et al., 2001;
Yamamoto et al., 1997). In keeping with their role as auto- and hetero-receptors, H3R
are heterogeneously distributed among areas known to receive histaminergic projec-
tions (Hu and Chen, 2017). High densities are found particularly in anterior parts
of the cerebral cortex, hippocampus, amygdala, nucleus accumbens, striatum, olfactory
tubercles, cerebellum, substantia nigra, and brain stem. In the TMN, H3R reside on
perikarya of histaminergic neurons. Loss of H3R function in KO mice is associated
with behavioral state abnormalities, reduced locomotion (Toyota et al., 2002), a met-
abolic syndrome with hyperphagia, late-onset obesity, increased insulin and leptin
levels (Tokita et al., 2006; Yoshimoto et al., 2006), and an increased severity of neu-
roinflammatory diseases (Teuscher et al., 2007).

1.2.4 Histamine H4 Receptor

Six independent laboratories contributed to the identification and cloning of the
H4 receptor (Liu et al., 2001; Morse et al., 2001; Nakamura et al., 2000; Nguyen et
al., 2001; O’Reilly et al., 2002; Oda et al., 2000; Zhu et al., 2001). The human H4
receptor gene is present on chromosome 18q11.2 and is a 44 kDa 390-amino-acid
polypeptide. Like H3R, the H4 receptor is coupled to pertussis toxin sensitive Gi/o
protein with inhibitory effect on cAMP accumulation (Leurs et al., 2009; Oda et al.,
2000). H4 receptor expression has been observed in eosinophils, T cells, dendritic
cells, basophils, and mast cells (Gantner et al., 2002; Hofstra et al., 2003; Liu et al.,
2001; O’Reilly et al., 2002), but its expression in the central nervous system remains
controversial. In the human brain, expression of H4 receptor mRNA has been reported
in the amygdala, cerebellum, corpus callosum, cortex frontal cortex, hippocampus,
and thalamus (Strakhova et al., 2009) but, results obtained with analyses of mRNA
expression does not always reflect results obtained with immunohistochemistry,
therefore there is a debate about H4R in SNC that needs further research (Panula et
al., 2015).

1.3 Homeostatic Histaminergic Functions

The morphology of brain histaminergic system with a compact group of cells and
capillary distribution of varicose fibres suggest its action as a normative centre for
the brain activity. Pharmacological studies in intact and histamine-deficient animals
as well as humans link brain histamine with homoeostatic brain functions and neuro-
endocrine control. Brain histamine controls behavioural responses, biological
rhythms, body weight, energy metabolism, thermoregulation, fluid balance, stress, and
reproduction (Hough, 1988; Parmentier et al., 2002; Schwartz et al., 1991).

Recently, our laboratory demonstrated functional differences in TMN neurons,
suggesting that histaminergic neurons are organized in distinct subpopulation imping-
ing on different brain regions (Blandina et al., 2012; Giannoni et al., 2009).
Histaminergic neurons help sustain wakefulness, several studies corroborate this hypothesis; in H1R-KO mice the sleep-wake cycle is impaired and the waking promotion induced by H1R antagonist is abolished (Huang et al., 2006; Lin et al., 2002). During waking c-fos expression increases in TMN neurons (Lin, 2000; Nelson et al., 2002; Nelson et al., 2003; Scammell et al., 2000; Sherin et al., 1998; Vanni-Mercier et al., 2003). The regulation of the transition between wakefulness and sleep involves antagonist influences of sleep-promoting VLPO neurons, which provide inhibitory GABA- and galanin-mediated inputs to TMN and brainstem cholinergic and monoaminergic groups, and excitatory effects of orexin (Hcrt/Orx) neurons on TMN and other wake-active neuronal groups (Benarroch, 2010). TMN neurons become active just after waking and fire at an average rate of about 5 Hz, and their activity is suppressed during sleep (Sakai et al., 2010; Saper et al., 2010; Takahashi et al., 2006). A recent elegant work by Wisden and co-workers demonstrated that zolpidem, a GABA_A receptor-positive modulator, needs to work on specific cell types of the brain, including histaminergic neurons, to induce sleep, without reducing the power of the sleep, hence improving sleep quality (Uygun et al., 2016). Furthermore, the same laboratory showed that wake-active histaminergic neurons generate a paracrine GABAergic signal that serves to provide a brake on over-activation from histamine, but could also increase the precision of neocortical processing (Yu et al., 2015b).

1.3.2 Thermoregulation

The key anatomical sites that control thermoregulation are under histaminergic innervation. The control centres of thermoregulation are located in the anterior preoptic area (thermoregulator rostral center) and in posterior hypothalamic area (thermoregulator caudal center) (Nieuwenhuys et al., 2008). Central administration of histamine in freely moving animals causes hyperthermia or biphasic responses, hypofollowed by hyperthermia (Clark et al., 1975; Clark and Cumby, 1976; Cote and Harrington, 1993). Hyperthermia, in turn, facilitates neuronal histamine release (Kanamaru et al., 2001). Activation of H1Rs and H2Rs in the anterior hypothalamic/preoptic area and posterior hypothalamus respectively induced hypothermia (Benarroch, 2010; Clark and Cumby, 1976). By H1Rs and H2Rs agonist actions, histamine regulates body temperature and clock neurons, suggesting an evolutionary conserved link between histamine, circadian rhythms, and temperature control.

1.3.3 Fluid Balance

Histamine elicits drinking following injection into the cerebral ventricles or into several hypothalamic sites (Gerald et al., 1972; Leibowitz, 1973). In addition, histamine increases the release of vasopressin and decreases urine output via both H1 and H2 receptors (Bennett and Pert, 1974; Bhargava et al., 1973; Kjaer et al., 1994). Histamine also stimulates vasopressin release indirectly via a local release of noradrenaline (Bealer, 1993; Bealer and Abell, 1995). A physiological role for central histamine in the control of fluid balance is suggested by the findings that 24 or 48 h of dehydration increases synthesis and release of histamine in the hypothalamus (Kjaer et al., 1994; Kjaer et al., 1995). Furthermore, blockade of histamine synthesis by α-FMH,
Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide activation of presynaptic H₃ autoreceptors or antagonism of postsynaptic histamine receptors all strongly depress dehydration-induced vasopressin release (Kjær et al., 1994). Dehydration induced renin release also appears to involve central histamine activation of the sympathetic nervous system (Kjær et al., 1998; Matzen et al., 1990).

### 1.3.4 Feeding and Energy Metabolism

The evidence for histamine involvement in food intake is nowadays very consistent (Provensi et al., 2016; Provensi et al., 2014). Early studies demonstrated that i.c.v injection of histamine, loading with histamine precursor L-histidine or application of the H₃ receptor antagonist thioperamide suppress feeding (Cohn et al., 1973; Machidori et al., 1992; Ookuma et al., 1993; Sheiner et al., 1985), whereas i.c.v infusions of a-FMH or H1 receptors antagonists increase food intake (Fukagawa et al., 1989; Ookuma et al., 1989; Sakata et al., 1988a; Sakata et al., 1988b). However, the role of histamine is not restricted to feeding control but also the regulation of body weight and adiposity by modulation of peripheral energy. Many of the central hypothalamic areas involved in regulating feeding, including the arcuate, ventromedial (VMH) and paraventricular (PVN) nucleus and lateral hypothalamic perifornical area (LHA), are densely innervated by histamine containing fibres and show a high density of H₁Rs (Panula et al., 1989). Early work suggested that histamine-mediated suppression of food intake was controlled by the VMH as microinfusion of H₁R antagonists into the VMH but not PVN or LH elicited feeding responses and increases both meal size and duration (Fukagawa et al., 1989; Sakata et al., 2003). Likewise, electrophoretic application of H₁R antagonists suppressed the firing of glucose-responsive units in the VMH but not in the LHA or PVN (Fukagawa et al., 1989). Another site of importance in the histamine control of food intake is the mesencephalic trigeminal nucleus. Bilateral injections of a-FMH into this region reduced eating speed and prolonged meal duration while leaving meal size unaltered. Feeding induced increases in histamine turnover in both the trigeminal nucleus, which controls mastication, and the ventromedial area, which is considered as a satiety center (Fujise et al., 1998). In our laboratory, we recently showed that the PVN as well takes part into the histaminergic control feeding behaviour as histamine released in the PVN activates oxytocin neurons (Provensi et al., 2014) that in turn exert hypophagic behaviour (Gaetani et al., 2010). Furthermore, the orexigenic actions of orexins/hypocretins (Jørgensen et al., 2005) and the anorexigenic effects of leptin (Toftegaard et al., 2003) and glucagon-like peptide-1 (GLP-1), which depend on CRH released by PVN neurons (Gotoh et al., 2005), are all blunted or absent by pharmacological or genetic loss of H₁R function. Ghrelin, another peptide of peripheral origin, does not affect histamine release, suggesting that ghrelin may act on a parallel, different mechanism that controls food intake (Ishizuka et al., 2006).

Most experimental observations in rodents agree that blockade of brain H₃ receptor, hence increasing histamine release, decreases energy intake, body weight and plasma triglycerides (Hancock and Brune, 2005). Also, they increase histamine release from the hypothalamus, they reduce energy intake in normal and leptin-resistant mice with diet induced obesity (Ishizuka et al., 2008), and decrease food intake in wild type mice (Provensi et al., 2014). From the pharmacological point of view the importance of histamine in the regulation of feeding behaviour came from the observation that increased weight is a common adverse effect of many classic antipsychotic
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drugs and atypical antipsychotics that depends on their affinity as antagonists at the H3Rs (Kroeze et al., 2003). Preclinical studies showed that activation of histamine neurons induces the arousal state during food anticipation (Angeles-Castellanos et al., 2004; Inzunza et al., 2000), and during the appetitive phase that precede food consumption (Passani and Blandina, 2011).
Chapter 2

Cognitive functions of brain histamine

The regulation of memory consolidation by histamine is a topic of much recent research and a new role for the brain histaminergic system is emerging, as a gateway between peripheral signals and the elaboration of both aversive and emotionally neutral memories (Passani et al., 2017).

Memory is not a unitary faculty of the mind but is a complex process composed and coordinated by multiple systems, distinct brain regions and pathways involves. Two memory systems have been described: declarative and procedural or 'habit' memory system (Izquierdo et al., 2006b). Declarative memory (“knowing that”) refers to conscious memory for events and facts, is assessed by explicit tests of recall and recognition, and depends upon medial temporal lobe and diencephalic brain structures. Procedural memory (“knowing how”) refers to unconscious memory, is assessed by experience-dependent learning of skilled performance, and depends on structures in the basal ganglia, cerebellum, and neocortex (Gabrieli, 1998). The memory systems of the mammalian brain operate independently and in parallel to support behaviour (McDonald and Hong, 2013; Packard and Goodman, 2013; Poldrack and Packard, 2003).

Procedural or working memory (WM) was considered as overlapping with immediate memory or short-term memory (STM). STM was assumed to be essential for acquisition and perhaps subsequent long-term memory (LTM) formation. It is generally accepted that the STM, that lasts minutes or few hours, may be converted in LTM that persists for days, weeks or more, by a specific sequence of events called consolidation which starts immediately after the acquisition phase (Izquierdo et al., 1999; McGaugh, 2000). Not all information reaching the CNS is stored; most inputs are filtered away by the attentional and emotional processes after the acquisition phase (Cahill and McGaugh, 1998). Only memories relevant for cognition, emotionally salient, or derived from strong sensory inputs are consolidated in LTMs. Forgetting means entails screening, since only memories previously selected persist, are stored and retrieved by chains of cause and effect controlled by a range of factors that operate simultaneously at many different levels often implicit, but sometimes explicit. The brain systems responsible for learning and storing memories have a mechanism for preventing information overload (Izquierdo et al., 2006b). Memory can be modulated by experiences occurring about the time when it is learned, consolidated or retrieved (Cahill and McGaugh, 1998; Kandel and Squire, 2000).
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2.1 Recognition Memory

Recognition memory involves at least two separable processes, familiarity discrimination and recollection and more complex aspects of contextual, associative and spatial aspects (Brown and Banks, 2015). Types of stimuli, single or multi-item, associative or non-associative stimuli, and types of information being stored, for example, the ‘what’, ‘when’, and ‘where’ of episodic-like memories or the modality of the sensory input allows the distinctions between types of memory process.

Discrimination for visual stimuli seems to be affected by a system centred on the perirhinal cortex of the temporal lobe (Brown and Banks, 2015). Regions other than perirhinal cortex may also be involved in recognition memory processes: when a recognition memory task is solved using recollection or association concerning a presented stimulus, the recognition memory predicts hippocampal involvement (Brown et al., 2010). The hippocampus is involved in recognition memory using multiple items and associative or spatial information (Aggleton and Brown, 2006; Dere et al., 2006; Eichenbaum et al., 2007; Murray and Ranganath, 2007; Squire et al., 2007; Winters et al., 2008). In particular, lesions of the rat hippocampus impair recognition memory that requires spatial information; location tasks are impaired by hippocampal lesions while perirhinal lesions have no effect (Warburton and Brown, 2010).

Functional imaging in human subjects has implicated the prefrontal cortex in recognition memory processes (O’Neil et al., 2012). The role of the rodent medial prefrontal cortex in recognition memory has been extensively studied. Large lesions of the prefrontal cortex, which included the anterior cingulate, prelimbic and infralimbic cortices, or which centred on the ventral medial prefrontal cortex, produced recognition impairments (Kolb et al., 1994; Ragozzino et al., 2002). The medial prefrontal cortex has been implicated in attentional processing (Chudasama and Robbins, 2003; Muir, 1996), and play an important role in temporal order memory (Chiba et al., 1997; Hannesson et al., 2004; Mitchell and Laiacona, 1998). Lesions in the medial prefrontal cortex impaired temporal order memory task (Barker et al., 2007; Devito and Eichenbaum, 2011) but not induces deficits in the recognition or location tasks (Barker et al., 2007). Moreover, in humans and non-human primates damage to the medial dorsal thalamus (MD nucleus) produces recognition memory deficits (Parker et al., 1997; Victor, 1987; Warburton and Brown, 2015).

2.2 Fear Memory

Forming associations about events and then consolidating memories of those associations is an important strategy for survival. However, in traumatic situations, these associations sometimes become overly consolidated and then, potentially, are resistant to extinction over time, resulting in fear-related disorders (Parsons and Ressler, 2013). But fear, in general, has a strong survival value. The lack of fear, also called recklessness or mindlessness in humans, is inherently dangerous and potentially lethal (Izquierdo et al., 2016). Therefore, in both cases, over-consolidation/resistant-extinction and recklessness/loss of fear are two side of the same coin, dangerous in the same way.

The acquisition and memory of conditioned fear depend on both hippocampus and amygdala, as lesion studies (Lorenzini et al., 1996a; Lorenzini et al., 1996b; Sacchetti et al., 1999; Sacchetti et al., 2002) and biochemical studies (Trifilieff et al.,

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2007) indicate. The sensory-related information from hippocampus and amygdala presumably originates in the mesencephalic reticular formation and ventral tegmental area (VTA), which receive it, in turn, from collaterals of the sensory pathways (BUSER and ROUGEUL, 1961; GREEN and MACHNE, 1955; MACHNE and SEGUNDO, 1956). The hippocampus and the basolateral amygdala (BLA) together with the ventro-medial prefrontal cortex (vmPFC) orchestrate memory formation (Izquierdo et al., 2016). Also the periaqueductal grey (PAG), is a brain region that conveys aversive signals to the amygdala. The PAG is known as an output structure for various conditioned fear responses, it receives a strong nociceptive input from the spinal and trigeminal dorsal horn (Gross and Canteras, 2012) and a recent study found that temporary pharmacological inactivation of PAG reduces shock-evoked responding in amygdala neurons and the acquisition of fear learning (Johansen et al., 2011).

2.3 How to evaluate memory in rodents: most widely used paradigms

Fear learning is usually studied by classical (Pavlovian) or instrumental association between the environment or changes in the environment (conditioned stimulus, CS) and a fearsome stimulus (usually one or more mild foot shocks; unconditioned stimulus, US). This type of learning represents situations in humans in which initially neutral stimuli become threatening through pairing with other stimuli and generate fear, a human emotion that guides much of our behavior and is crucial for survival (Izquierdo et al., 2016).

In the fear conditioning paradigm, the animals are placed in a new environment (context) were they receive a mild aversive stimulus such as a foot shock (US) associated with another stimulus such as tone or light (CS), that usually does not elicit a response. Following learning, the presentation of the CS alone generates various visceral and behavioural conditioned fear responses. The term fear response is used to refer specifically to measurable responses that occur in response to threat and not to the conscious feelings of fear: called freezing behaviour. Freezing behaviour (conditioned response, CR), is a generalized immobility caused by a generalized tonic response of the animals' skeletal musculature except those muscles used in breathing (Herry and Johansen, 2014; Izquierdo et al., 2016).

Promnesic agents are expected to increase, whereas amnesic manipulations to reduce freezing behaviour (Curzon et al., 2009; Wehner and Radcliffe, 2004).

The most widely used instrumental fear conditioning is one-trial inhibitory avoidance (IA) (Gold, 1986; Izquierdo et al., 2006a; Izquierdo and Medina, 1997) which used to be called “passive avoidance” in opposition to the “active avoidance” tasks in which animals have to perform some movement to avoid the foot shocks. In the “passive” tasks animals have to withhold stepping through a hole into a dark compartment, or stepping down from a platform onto a grid, to access the shock compartment; the required response is to remain in the safe, lit compartment or on the start platform. Animals learn to avoid stepping through or stepping down, but they are not in any way refrained from moving or behave in any way passively. In fact, they move a lot while on the platform or in the lit compartment (Netto and Izquierdo, 1985). When retested, an increase in the latency to step-through or to step-down is related as a measure of learning (Izquierdo and McGaugh, 2000).

During the last decade the ‘what’, ‘where’ and ‘when’ (WWWhen) episodic-like memory (ELM) task, which is based on the object recognition paradigm, has been
Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide utilized for the cognitive phenotyping of mouse mutants and transgenic mouse models of neuropsychiatric diseases. It was also widely used to identify the neuroanatomical, electrophysiological and pharmacological foundations of ELM formation, retention and retrieval (Binder et al., 2015). The object recognition paradigm is one-trial task, does not involve learning of rules, does not require reinforces and is purely based on the innate preference of the rodents to explore the novel object. Thus, a rodent that remembers the familiar object will spend more time exploring a novel object rather than the familiar one (Leger et al., 2013). The preference for the novel object is the result of incidental learning occurring during the sample phase and exploratory behaviour during the testing phase (Jablonski et al., 2013).

2.4 The role of central histaminergic system in memory and cognition

Recent findings have reasserted the role of histamine in the regulation of memory consolidation first proposed in 1986 in an inhibitory avoidance task in rats (Passani et al., 2017). The first description on the role of histamine in memory process came from de Almeida and Izquierdo, 1986 (de Almeida and Izquierdo, 1986); the authors demonstrated that the infusion of histamine in the lateral ventricles facilitates fear memory process when tested in the inhibitory avoidance paradigm. At that time, though, it was not entirely clear, however, whether histamine had a physiological role in memory modulation. We now know that histamine exerts different effects in different brain regions and different modulatory actions depending on the type of memory task and the phase of memorization (Passani et al., 2017; Passani et al., 2007). In the hippocampus and amygdala the histaminergic neurotransmission exerts different actions. Histamine or H3 antagonists infusion in the hippocampus ameliorated the performances in spatial learning (Huang et al., 2003). Histamine infusion in the dorsal hippocampus facilitates fear memory by activating H2 receptor (da Silva et al., 2006), whereas in the ventral hippocampus H1 receptor inhibits fear memory (Alvarez and Banzan, 2008).

When histamine was locally injected into dorsal and ventral regions of the hippocampus improvements in the rats performances in the inhibitory avoidance test (Alvarez and Banzan, 1996, 2001, 2008; da Silva et al., 2006) and also in the radial-maze tasks (Huang et al., 2003; Yamamoto et al., 2007) were observed. Conversely, blockade of endogenous histamine production by α-FMH infusion into the ventricles inhibits the consolidation of IA, which is known to be sustained by both the BLA and the hippocampus (Izquierdo et al., 2006a; Izquierdo et al., 1992; Izquierdo and Medina, 1997; McGaugh, 2000, 2015).

The H1 and H2 histamine receptors in BLA, hippocampus, and vmPFC facilitate memory consolidation in the inhibitory avoidance task (IA) and contextual fear conditioning (CFC), and their specific antagonists have an opposite effect in the consolidation of different tasks (Benetti et al., 2012; Benetti and Izquierdo, 2013; Fiorenza et al., 2012).

A recent, seminal observation by Benetti et al. (2015) (Benetti et al., 2015) revealed that histaminergic neurotransmission provides the brain with the plasticity necessary to ensure memorization of emotionally salient events, through recruitment of alternative circuits. The authors found that the integrity of the brain histaminergic system is necessary for long-term, but not for short-term memory of step-down inhibitory avoidance. In addition, they observed that phosphorylation of cyclic adenosine
monophosphate responsive-element-binding protein (CREB), a crucial mediator in long-term memory formation (Bernabeu et al., 1997; Josselyn et al., 2004) correlates anatomically and temporally with histamine-induced memory retrieval, showing the active involvement of histamine in the CA1 region of the hippocampus and in the basolateral amygdala in different phases of memory consolidation.
Chapter 3
Disorders associated with brain histamine
No disease entity has so far been linked specifically or selectively to brain histamine dysfunction, but histamine dysfunction may thus be a precipitating factor for epigenetic disease susceptibility, severity, and progression.

Histamine as the major wake-promoting neurotransmitter in the CNS and plays a role in the pathogenesis of sleep disorders. Cerebrospinal fluid histamine levels are decreased in people with idiopathic hypersomnia or narcolepsy, supporting the concept that histamine in humans promotes alertness and wakefulness (Kanbayashi et al., 2009; Nishino et al., 2009). Narcolepsy is a sleep disorder characterized by excessive daytime sleepiness, catalepsy and narcoleptic episodes. Insomnia is usually treated with benzodiazepines, whereas narcolepsy was treated until recently with wakefulness-promoting compounds, like modafinil, and amphetamines that modulate the dopaminergic system. In 2015, the European Medicines Agency (EMA) has recommended pitolisant, a first-in-class medicine that acts on histamine H3 receptors in the brain that blocks autoreceptors, for the treatment of narcolepsy with or without cataplexy.

In contrast to other aminergic systems, both histamine and the levels of its metabolites increase in the spinal fluid with increasing age (Prell et al., 1988). In Alzheimer’s disease, several subcortical ascending projections, including the histaminergic neurons, display degeneration and tangle formation (Swaab et al., 1998). In the hypothalamus, neurofibrillary tangles occur exclusively in the TM nucleus (Airaksinen et al., 1991b) and histamine levels is reduced in different brain areas including hippocampus, hypothalamus, frontal and temporal cortex (Panula et al., 1998; Shan et al., 2012) suggesting that histaminergic neurons undergo degeneration and contribute to cognitive decline in this disorder. H3R antagonists improve cognitive performance in experimental animals. So far, though, clinical trials with these compounds have proven unsuccessful (NIH website). The TM neurons seem morphologically normal in patients with Parkinson’s disease (PD) (Nakamura et al., 1996), and normal histidine decarboxylase activity has been observed (Garbarg et al., 1983). In contrast, histamine levels are markedly increased in the substantia nigra, putamen and globus pallidus (Rinne et al., 2002), but only modest changes in t-MeHA, the main metabolite of histamine, and histamine-N-methyltransferase activity (Fogel et al., 1994) indicating that the increased histamine might not be present in the releasable pool in the brains of these patients. This might be due to the limited capacity of the vesicular monoamine transporter VMAT-2 to take up histamine into the vesicle.
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Thus, the alteration in histamine may be a direct consequence of dopamine deficiency, a hallmark of Parkinson’s disease. The abnormalities in the histaminergic system is highlighted in PD patients which often experienced fragmented sleep and insomnia. Alteration in the histaminergic system is also revealed in receptor binding profile. H₃R binding is abnormally high in the Parkinsonian substantia nigra (Anichtchik et al., 2000), and the same phenomenon is seen in rats after depletion of nigrostriatal dopamine stores using 6-OHDA (Ryu et al., 1994). However, H₃R antagonists improve motor coordination in neonatally 6-OHDA-lesioned rats, whereas imetit, an H₁R agonist, attenuates 3,4-dihydroxyphenylalanine (l-DOPA)-induced increases in striatal extracellular dopamine and dyskinesia (Nowak et al., 2008). Histamine, via activation of H₃Rs, thus regulates the cortico–striato–nigral system at all levels (Panula and Nuutinen, 2013).

The first antipsychotics were developed as antihistaminic compounds, before dopamine was recognized as a key neurotransmitter in the brain. It is now known that the atypical antipsychotic drugs have an affinity for many different receptors, including those for serotonin, noradrenaline and histamine. In the brains of people with schizophrenia, various changes are found in the histaminergic system (Jin et al., 2009; Prell et al., 1995). Levels of t-MeHA are elevated in the spinal fluid of patients with schizophrenia (Prell et al., 1995), this implies that there is increased histamine release and turnover that could explain the low H₁-receptor binding that is observed in the frontal cortex of people with schizophrenia (Nakai et al., 1991). Histamine–H₃R binding is increased in the dorsolateral prefrontal cortex but unchanged in the temporal cortex of patients with schizophrenia compared with those brain regions in healthy control subjects (Jin et al., 2009). These differences may reflect the structural abnormalities of the cortical network and changes in cellular composition that underlie the functional impairments in this disorder. All antipsychotics act on dopamine D₂R, supporting the proposition of dopaminergic supersensitivity as a major factor in disease susceptibility and pathogenesis (Seeman et al., 2006). H₃R antagonist/inverse agonist increase the dopamine and acetylcholine levels in the cortex but not in striatum, suggesting that negative symptoms and cognitive deficits could be ameliorated. Preclinical and clinical studies indicate that H₃R antagonist/inverse agonist are not sufficient as antipsychotic therapy per se but represent a promising poly-therapy treatments in schizophrenics. Thus, antagonist/inverse agonist H₃R, D₂R, 5-HT₂A combination, could be considered a rational approach in different types of psychosis treatments (Ito, 2009; Tiligada et al., 2009; von Coburg et al., 2009).
Chapter 4
Is histaminergic neurotransmission involved in antidepressant responses?

4.1 Historical background

The term depression began to appear in the 19th century as did the modern concept of affective disorders, with the core disturbance now viewed as one of mood. Melancholia was recognized as early as the time of Hippocrates and continued through galenic medicine and medieval times. The earlier connotation of the term was very wide and included all forms of quiet insanity. Melancholia later became more clearly associated with the more modern idea of melancholy or despair, for instance, in the classic work of the English Renaissance author, Richard Burton, The Anatomy of Melancholy, first published in 1621.

The first division of the major endogenous psychoses is related to Kraepelinian dichotomy that delineates the foundations of the modern classification of psychiatric disorders. This division was formally introduced in the sixth edition of Emil Kraepelin's psychiatric textbook Psychiatrie. Ein Lehrbuch für Studirende und Aertze, published in 1899 (Decker, 2007). Kraepelin regarded psychiatric disorders as disease entities based on a medical, neurological model, with specific, organic etiology and pathology. He did, however, regard some pathological depressions as psychogenic in origin. While he did not completely clarify his views on their position in his classification, or how they were to be distinguished from manic-depressive illness with incidental stress, he appeared to regard them as a separate, but relatively small and unimportant group.

Another school of European psychiatrists was developing a very different approach. The psychoanalysts. Freud and Abraham developed a theory of the origin of depression in relation to actual or symbolic losses of a love object. Here was a theory regarding the origin of most, if not all, depressions as psychogenic. Psychological theories of causation became more widely accepted for these disorders. A challenge now arose as to how to reconcile these theories with older ones of organic causation. Adolf Meyer, a Swiss psychiatrist moved away from the idea of clear cut disease entities, and viewed all psychiatric disorders as reaction types, or psychobiological reactions of the organism to stress (Meyer, 1922). The Meyerian concept defined depression as a type of reaction in which both psychological and organic factors had to be taken into account (Paykel, 2008).
4.2 The modern concept of depression

Depression is among the leading contributors to the global burden of disease (Whiteford et al., 2013), globally, more than 300 million people of all ages suffer from depression. It is associated with enormous personal suffering and societal economic burden (Kessler et al., 2003) with 10-30% of women and 7-15% of men likely to suffer from depression in their life-time (Briley and Moret, 2000). Furthermore, depression can be a lethal illness owing to an elevated risk for suicide (Trivedi et al., 2006), as well as increased risks of cardiac disease, cerebrovascular disorders and other medical causes of mortality (Walker et al., 2015). The magnitude of the clinical burden of depression reflects, in part, the limited effectiveness of present-day treatments. Currently available antidepressant medications, alone and in combination, are associated with high rates of partial responsiveness or non-responsiveness, delayed response onset of weeks to months and limited duration of efficacy (Gaynes et al., 2012). Despite the high prevalence of depression and its socioeconomic impact, the etiology and pathophysiology of this complex disorder is not well understood. It is the lack of understanding of the underpinning of depression that has resulted in no substantial improvement to antidepressant treatment (Misra, 2012) and consequently the lack of improvement over the conventional monoaminergic-based therapies discovered by serendipity decades ago (Covington et al., 2010; Kessler et al., 2008).

A major liability of the monoamine-deficiency hypothesis is its derivation from the mechanism of currently available antidepressants. Despite their efficacy, however, current antidepressant pharmacotherapy alleviates symptoms in approximately two thirds of patients proving a clinical response to these agents, whereas one third have a response to placebo (Mann, 2005). Perhaps the mechanism of depression is not related to monoamines in two of three cases (Belmaker and Agam, 2008). Indeed, the monoaminergic theory for depression is not able to explain the fact that biochemical effects of antidepressant rapidly occur whereas the therapeutic response has a delay of days or weeks (Stahl, 2008). Thus, the immediate effects of antidepressant drugs on monoamines cannot fully explain this lag period of treatment response (Malberg and Blendy, 2005).

To better understand mechanisms underlying the therapeutic efficacy of antidepressant drugs, research efforts have focused on the long-term molecular changes that underlie depression and antidepressant treatments.

4.3 Neurogenic and neurotrophic theory

The supposition that monoamine deficits may not reflect a core feature of depression pathophysiology, but are the result of neural dysfunction directed research away from monoamines and towards the putative role of growth factors such as brain-derived neurotrophic factor (BDNF), known to be critically involved in regulating neural structure and plasticity in the adult brain (Kafitz et al., 1999; Thoenen, 1995). The first studies to implicate BDNF in antidepressant responses showed that conventional antidepressant drugs, as well as electroconvulsive therapy (ECT), enhanced BDNF and TrkB mRNA expression in the hippocampus and cortical regions in a timeframe similar to the onset of antidepressant-like response (Nibuya et al., 1995; Nibuya et al., 1996). To more directly examine the causal involvement of BDNF in antidepressant
responses, Siuciak and colleagues infused BDNF protein directly into the midbrain and observed an antidepressant-like effect in rodents (Siuciak et al., 1997).

In humans, acute and chronic stress both decrease endogenous neurotrophin levels and can lead to significant atrophy of the hippocampus, a structure known to be involved in controlling emotionality (Duman, 2004; McEwen, 2000; Sapolsky, 1996). These events may be causally linked via neurogenesis (Duman, 2005).

Neurotrophins are growth factors with crucial roles in the formation and plasticity of neuronal networks (Huang and Reichardt, 2003). The neurotrophin family include nerve growth factor (NGF), BDNF, neurotrophin-3 (NT-3) and neurotrophin-4 (NT-4). They are initially synthesized as precursor proteins (pro-neurotrophins), which are processed extracellularly or intracellularly to be secreted mostly in a mature, biologically active form (Matsumoto et al., 2008; Yang et al., 2009). The neurotrophins, show binding specificity for particular tyrosine kinase receptors (Trk), and non-specificity for the p75 neurotrophin receptor (p75NTR). In particular, pro-BDNF preferentially activates p75NTR receptor to mediate programmed neuronal death (Frade et al., 1997), to decrease dendrite complexity and spine density in hippocampal neurons (Zagrebsky et al., 2005), and to induce long-term depression of synaptic transmission (Rösch et al., 2005; Woo et al., 2005). On the contrary, mature BDNF (mBDNF) selectively activates TrkB, a member of the tyrosine kinase receptors family, to promote survival and differentiation, increasing the branching of axons and dendrites and stabilizing synaptic contacts (Lee et al., 2001). Thus, due to the essential role of BDNF for cell differentiation, nerve growth and neuronal survival it has been implicated in several brain diseases, including depression considering its well establishes relation with low levels of BDNF expression (Neto et al., 2011).

Birth of new neurons or neurogenesis continues to occur in selected neurogenic zones in the adult brain. This includes the subventricular zone that gives rise to olfactory bulb neurons, and the subgranular zone that generates granule cells of the hippocampal dentate gyrus. Similar to regulation of BDNF in the dentate gyrus, stress and antidepressant treatments exert opposing effects on neurogenesis in the adult hippocampus. Stress and depression decrease the expression and function of BDNF in the PFC and hippocampus, structures that are implicated in depression, as well as decrease the BDNF levels in the blood of subjects with depression (Bocchio-Chiavetto et al., 2010; Krishnan and Nestler, 2008; Turner et al., 2006). Reduced neurotrophic or growth factor levels may be particularly relevant to the structural alterations caused by stress and depression, as these factors (particularly BDNF) are required for activity-dependent formation and maintenance of synaptic connections (Holtmaat and Svoboda, 2009; Joudi et al., 2009). Studies of a human BDNF polymorphism (BDNFVal66Met) that is found in approximately 25% of the population have been insightful. The presence of the BDNFVal66Met allele, whose product results blocks the processing and release of mature BDNF, is sufficient to cause atrophy of neurons in the hippocampus (Chen et al., 2006) and medial PFC (mPFC) of mice with this allele (Liu et al., 2012). Heterozygous deletion of BDNF also decreases spine density and dendrite length of hippocampal and PFC neurons, decreases hippocampal volume and occludes the effects of chronic stress (Liu et al., 2012; Magariños et al., 2011). These findings suggest that stress could cause neuronal atrophy via inhibition of BDNF or that BDNF is required for neuronal remodelling (Duman et al., 2016).

A different point of view actually discusses the balance between neuronal death and neurogenesis as an etiological cause of this pathology (Jacobs et al., 2000;
Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide (Malberg et al., 2000). In fact, neurogenesis is a possible phenomenon in adults (Duman et al., 2001; Gould and Tanapat, 1999) and neurotrophins are important for nervous cells formation and differentiation (Fariñas et al., 2001; Fritzsch, 2003). Several studies have been proposed to test some of these theories relating the balance between neuronal death and neurogenesis, but most of them still lack experimental verification (Merighi et al., 2008).

4.4 Role of CREB in Depression and Antidepressant Treatments

The neurotrophic theory of depression led to a novel therapeutic approach in the management of depression beyond manipulation of neurotransmitter-receptor interaction, consisting in targeting signal transduction and gene expression pathways. The most investigated pathway, in that sense is the Cyclic AMP (cAMP) second messenger system, which ultimately influence gene expression by activating the transcription factor cAMP response element binding protein (CREB).

CREB is a nuclear protein. It belongs to the family of leucine zipper transcription factors that are expressed in a variety of tissues and serve diverse functions. Phosphorylation of a serine residue (S133) in its kinase inducible domain is critical to mediate its effect, as this permits recruitment of co-activator proteins and initiation of transcription (Nair and Vaidya, 2006). Activation of CREB can be accomplished by phosphorylation via cAMP-protein kinase A (PKA) pathway. PKA cascade also serves as a target for antidepressant treatment (Figure 3) (Tardito et al., 2006). CREB is regulated by diverse signalling pathways and integrates the action of numerous external stimuli, including antidepressant. A downstream consequence of enhanced CREB function is thought to be the increased expression of target genes like the neurotrophin, brain-derived neurotrophic factor (BDNF) (Figure 3), and neuropeptide Y (NPY), which may contribute to the antidepressant treatment mediated changes in structural plasticity and behaviour (Conti et al., 2002; Pandey, 2003). As previously mentioned, the hippocampus is a key limbic region whose structure and function are compromised in mood disorders. Hippocampal over-expression of CREB and BDNF can mimic both the structural consequences of sustained antidepressant treatment as well as exerting antidepressant like behaviour (Chen et al., 2001). Elevated CREB- BDNF, through their protective influences on vulnerable hippocampal neurons and their ability to directly promote structural recognition, could result in repair of damaged region due to depression (Misra, 2012). Elevated CREB function can either reduce or produce depressive-like behaviour in laboratory animals. For instance, elevated CREB activity in the NAc produces various depressive-like effects in rodents (Barrot et al., 2002). Until recently, the NAC has not been considered a likely site for the pathophysiology of depression, although it makes intuitive sense that symptoms of anhedonia, reduced energy and reduced motivation, which are prominent in many depressed patients, involve this brain reward region (Nestler et al., 2002; Pliakas et al., 2001). Within the amygdala, the consequences of alterations in CREB function appear to be state-dependent. Virus-mediated expression of CREB in the amygdala before training in the learned helplessness paradigm causes depressive-like effect, whereas expression after training results in antidepressant-like effect (Wallace et al., 2004). These finding provide further evidence that the actions of CREB are regionally and temporally specific. Therefore, the increased CREB function within the hippocampus produces antidepressant-like effects that correlate with elevated expression of...
growth factors such as BDNF, whereas the same increases in CREB function within the NAc produce many depressive-like signs. Such observations highlight the fact that CREB functions generally regulate plasticity, a process that is not good or bed; it could be adaptive, maladaptive, or both simultaneously. In the case of depression, elevations in CREB activity in one region and reduction in another could detract from the therapeutic action of treatment regimens that produce global influences on CREB function in the brain (Carlezon et al., 2005).

4.5 Neurocircuitry of depression

Regions within the orbital PFC (oPFC) and the mPFC appear to work as a coordinated unit to integrate sensory information, provide emotional salience and modulate visceral motor reactions and value-based decision processes (Wallis, 2011). These regions have connections with several sensory areas (Carmichael and Price, 1995), as well as inputs from the hypothalamus, amygdala, NAc and hippocampus. In rodents, the infralimbic PFC (IL-PFC) is believed to have roles similar to those of the oPFC-mPFC networks, by integrating information and modulating visceral reactions that are related to emotional processes through connections with the amygdala, hypothalamus and various brain stem nuclei (Vertes, 2004). Recent work suggests that the IL-PFC also modulates activation of the ventral tegmental area through effects on the amygdala and ventral subiculum, tying the region to subcortical reward-processing networks (Patton et al., 2013). Anhedonia, especially deficits in non-consummatory reward behaviour, is another core symptom of depression. Abnormal activity levels in the PFC-ACC, as well as in the ventral and dorsal striatum, have been reported in depressed patients with anhedonia (Pizzagalli, 2014).

Depression is also highly heritable, with roughly 40-50% of the risk for depression being genetic, although the specific genes that underlie this risk have not yet been identified. The remaining 50-60% of the non-genetic risk also remain poorly defined, with suggestion that early childhood trauma, emotional stress, physical illness, and even viral infections might be involved. Therefore, the highly variable compilation of symptoms that is used to define depression, and the highly variable course of the illness and its response to various treatments, indicate that depression subsumes numerous disease states of distinct etiology, and perhaps distinct pathophysiology. In fact, the lack of bona fide objective diagnostic test for depression, beyond a compilation of symptoms, means that the diagnosis of the syndrome is quite variable, with no clear line distinguishing people who have mild clinical depression from those who are simply having a tough time in the course of normal life (Berton and Nestler, 2006).

4.6 Animal Models used to screen antidepressant compounds

Despite the prevalence of depression and its serious impacts, studies on the pathogenesis of depression are still preliminary compared to those on the pathogenesis of other common chronic and potentially fatal multi-factorial conditions (Yang et al., 2010); the major obstacle is the restricted availability of validated animal models.

Ideally, an appropriate animal model of human depression should fulfill the following criteria as much as possible: strong phenomenological similarities and similar pathophysiology (face validity), comparable etiology (construct validity), and common treatment (predictive validity) (Vollmayr et al., 2007; Willner and Mitchell,
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Unfortunately, depression is a heterogeneous disorder and its many core symptoms (e.g., depressed mood, feeling of worthlessness, and recurring thoughts of death or suicide) are hard to be mimicked in laboratory animals.

It should be noted that there is a difference between a model and a test. A model can be defined as an organism (non-human) or a particular state of an organism that reproduces aspects of human pathology, providing a certain degree of predictive validity. A test, on the other hand, provides only an end-point behavioural or physiological measure (read-out) designed to assess the effect of a genetic, pharmacological or environmental manipulation (Urani et al., 2005).

While symptoms of depression are likely to be purely human features, other aspects of the depressive syndrome have been replicated in laboratory animals, and in several instances ameliorated with antidepressant treatment. These include measures of helplessness, anhedonia, behavioural despair and other neuro-vegetative changes such as alterations in sleep and appetite patterns (Krishnan and Nestler, 2011). Many of the test commonly used to detect antidepressant compound are essentially validated retrospectively based on the effects seen with clinically effective antidepressant agents, it is perhaps more appropriate to consider them as a test of antidepressant action rather than test of depression (Cryan and Mombereau, 2004).

I will mention only the most widely used preclinical tests used to screen antidepressant compounds.

4.6.1 Tail Suspension Test (TST)

The TST is based on the observation that rodents after initial escape-oriented movements, develop an immobile posture when placed in an escapable stressful situation, that in TST involves haemodynamic stress of being hung in an uncontrollable fashion by their tail (Thierry et al., 1986). If antidepressant treatments are given prior to the test, the subjects will actively persist engaging in escape-directed behaviours for longer periods of time than after vehicle treatment. The test is usually quite short, 6 minutes, and the amount of time they spend immobile is recorded (Steru et al., 1985). Similar to the FST, its validity is questioned by the fact that acute antidepressant treatments reverse the behavioural “depression” (Cryan and Mombereau, 2004). Although both the FST and TST are similar in the construct that they purport to asses, they are probably different in term of the biological substrates that underlie the observed behaviour although they often offer converging data on potential antidepressant (Bai et al., 2001; Porsolt, 2000; Renard et al., 2003).

4.6.2 Forced Swim Test (FST)

The Porsolt test (Porsolt et al., 1977), also known as the FST test is the most widely and most frequently used experimental paradigm to detecting antidepressant activity, largely due to its relative reliability across laboratories and its ability to detect activity in a broad spectrum of clinically effective antidepressant (Cryan et al., 2002; Porsolt, 2000). The test is based on the observation that rodents, following initial escape-oriented movements, develop an immobile posture in an inescapable cylinder filled with water. If antidepressant treatments are given prior to the test, the subjects will actively persist engaging in escape-directed behaviours for longer periods of time than after vehicle treatment (Cryan and Mombereau, 2004).
An interpretation of these tests as a model of depression is that immobility time is a symptom of reduced reactivity to an aversive environment, and the fact that the administration of an antidepressant prolongs the struggling or swimming time should give predictive validity to the model. A main criticism of this interpretation is that a very short-term treatment (acute or short-term 3-4 administration within 24h) is sufficient for shortening immobility time, and this fact conflicts with the delay necessary for an antidepressant compound to develop its therapeutic activity.

There are several differences between the two tests, such as their differential sensitivity to the immobility-reducing effect of various antidepressant with an apparent increased sensitivity of the TST. The mouse FST has not traditionally been viewed as a consistently sensitive model for detecting SSRIs activity (Porsolt and Lenegre, 1992) whereas these antidepressant are generally reported as active in the TST (Cryan et al., 2005).

Despite the major disadvantage of the TST related to its sensitivity to short-term antidepressant, the validity of the TST is based on the behavioural responses, comprising an evolutionary preserved coping strategy in which immobility behaviour represent the psychological concept of “entrapment” described in clinical depression (Dixon et al., 1998; Lucki, 2001). It is reasonably analogous to the human disorder in its manifestation or symptomatology, there are behavioural changes that can be objectively monitored, the behavioural changes observed should be reversed by the same treatment modalities that are effective in humans (Cryan et al., 2005). This fact would imply that TST should be considered a good model of antidepressant action.

4.6.3 Chronic Mild Stress (CMS)

One of the most elegant long-term models of depression is the chronic mild stress procedure devised by Willner (Papp et al., 1996; Papp et al., 1991). As the name suggests this paradigm consists of exposing rodents to a series of mild unpredictable stressors during a prolonged period (usually >2 weeks). This stress regimen induces many long-term behavioural, neurochemical, neuroimmune and neuroendocrine alterations resembling dysfunctions observed in depressed patients (Willner, 1997). In the CMS model, chronic sequential exposure to a variety of mild stressors has been shown to decrease the drinking of a sweetened solution, a condition that could be reversed by the chronic administration of classical antidepressant drugs as well as dopaminergic agonists (Muscat et al., 1992; Muscat et al., 1990; Papp et al., 1996; Willner et al., 1992). Exposure to chronic mild stress also impairs the acquisition of place preference conditioning, in parallel with sucrose consumption (Papp et al., 1991). These anhedonia-like behaviours have generally been shown to be reversed by chronic, but not acute, treatment with several classes of antidepressants (Moreau et al., 1992; Willner, 1997). Although the paradigm has been described as a model with a high predictive, construct and etiological validity (Cryan and Mombereau, 2004; Willner, 1990). Nevertheless, reservations concerning the reproducibility of the behaviour results obtained have been raised, thus questioning the reliability of the model.
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4.6.4 Learned Helplessness

This paradigm was originally developed based on the observations that dogs subjected to repeated inescapable uncontrollable (but not those subjected to controllable) shocks demonstrate escape deficits (Overmier and Seligman, 1967; Seligman and Maier, 1967). The model was later translated to the rat (Seligman and Beagley, 1975) and subsequently to the mouse (Anisman et al., 1979). The rodent studies revealed that the behavioural deficits are sensitive to a broad spectrum of antidepressants usually after short-term treatment (Leshner et al., 1979; Martin et al., 1990a; Martin et al., 1990b; Petty and Sherman, 1979). The major drawbacks of the learned helplessness model are two-fold: first most of the depression-like symptomatology does not persist beyond 2–3 days following cessation of the uncontrollable shock (Weiss and Kilts, 1998). Further, only a certain percentage (estimates vary somewhere between 10 and 80%) of animals develop helplessness behaviour (Drugan et al., 1989).

4.6.5 Novelty-suppressed feeding and sucrose preference

Novelty-suppressed feeding (NSF) test is a hyponeophagia-based behavioural test and provides an anxiety-related measure that is sensitive to the effects of chronic treatment with AD. The NSF also exhibits considerable potential as animal models for studying the neurobiology of the AD response (Dulawa and Hen, 2005). The NSF elicits competing motivations: the drive to eat and the fear of venturing into the centre of a brightly lit arena. Chronic, but not acute treatment with AD could significantly decrease the animal’s latency to eat, while it does not affect the food intake of animals in their home cage (Santarelli et al., 2003).

The sucrose preference test is usually used for analysing the effect of drugs on depressive animals. Reduced preference for sweet solution in sucrose preference test represents a loss of interest, fatigue and a loss of energy during depressive episodes, while this reduction can be reversed by treatment with ADs. The reduced sucrose preference has been used as a measure of anhedonia in the animal models of CMS and learned helplessness (Vollmayr et al., 2004; Willner et al., 1987).

4.6.6 Neuronal histamine: an insight on depression

Pharmacological or genetic loss of histamine or histamine receptor function in animals produces phenotypes that model human depression (Dai et al., 2007; Ito et al., 1999; Nath et al., 1988; Song et al., 1996). Histamine neurons in the TMN are sensitive to many, if not all, neuroendocrine signals implicated with depression, including biogenic amines, peptides, and steroid hormones, as well as antidepressant medication. Histamine neurons are strongly excited through 5-HT2C, a serotonin receptor that undergoes posttranscriptional editing (Sergeeva et al., 2007) that correlates with suicide (Schmauss, 2003). Noradrenergic α2-receptors increase GABAergic inhibition of TMN neurons (Nelson et al., 2003; Stevens et al., 2004), and interactions with peptidergic influences, e.g., hypocretins (Eriksson et al., 2001; Eriksson et al., 2004), CRH, and steroid hormones, may be implicated in neuroendocrine and coping abnormalities in depression.

PET studies using [11C]doxepin, an antidepressant with high affinity to H1R, revealed reduced H1R binding in frontal and prefrontal cortices, and the cingulate gyrus
correlating with the severity of clinical depression (Kano et al., 2004; Yanai and Tashiro, 2007). Anomalies in histamine metabolism (methylation) may account for endogenous depression in humans (Gagne et al., 1982), and the association of depression and atopy (Timonen et al., 2003) is in line with convergent roles of histamine in immune and stress responses (Steinman, 2004; Theoharides and Konstantinidou, 2007).

Munari and coworkers (2015) demonstrated that an intact histaminergic neurotransmission is essential for the SSRI citalopram and paroxetine to exert their behavioural and neurochemical actions in the TST paradigm. Munari et al., help to clarify also the putative interactions between the serotonergic and histaminergic systems in the antidepressant action of this two compound suggesting that SSRIs increase extracellular levels of endogenous 5-HT in the TMN, which in turn impacts 5-HT2C receptors localized on HA neurons, enhances their firing rate, and consequently augments HA release in the cortex. Disruption of this loop in HA-deprived mice is at least in part responsible for the inefficacy of citalopram and paroxetine (Munari et al., 2015).

Single nucleotide polymorphism of the HA H1R gene was found to play a role in bipolar disorder, as it was significantly associated with improvements following olanzapine and fluoxetine treatment (Perlis et al., 2010). In addition, functional mutation in the HDC gene resulting in deficits of the histaminergic neuronal system has been linked to the mechanism and modulation of Tourette’s syndrome and tics (Baldan et al., 2014; Ercan-Sencicek et al., 2010). Similar genetic variations in the population may contribute to individual differences in antidepressant response and may prove good predictors of more effective treatments (Munari et al., 2015). Modulation of histaminergic system may thus prove to be useful in the treatments of depression and related mood disorders.
Chapter 5

Is neuronal histamine involved in stress-related responses?

5.1 Historical Background

Aristotle, Hippocrates, and the other Ancients were aware of stress and its adverse effects. However, Claude Bernard was the first to formally explain how cells and tissues in multicelled organisms might be protected from stress. Bernard, first pointed out (1859) that the internal medium of the living organism is not merely a vehicle for carrying nourishment to cells. Rather, is the condition of life. Fifty years later, Cannon, working at Harvard, suggested the designation homeostasis for the coordinated physiological processes that maintain most of the steady states in the organism. Cannon coined the term 'fight or flight' to describe an animal's response to threat. The concept of 'fight or flight' proposes that animals react to threats with a general discharge of the sympathetic nervous system, priming the animal for fighting or fleeing. This response was later recognized as the first stage (acute stress response) of a general adaptation syndrome (GAS) postulated by Hans Selye to be a universal stress response among vertebrates and other organisms. Selye, also known as the "father of stress", observed that patients with a variety of illnesses had many of the same "non-specific" symptoms that were a common response to stressful stimuli experienced by the body. These clinical observations together with experiments on laboratory rats underpinned Selye's concept of GAS, which led Selye to assert that prolonged exposure to stress resulted in "diseases of adaptation". That is, chronic stress, by causing the overproduction of chemicals and hormones, produced gastroduodenal ulcers and high blood pressure. Although the GAS hypothesis was subsequently shown to be incorrect, it did put stress on the map and also highlighted the fact that stress had major effects on the immune system as well as on the adrenal glands. In addition to providing the first clear definition of stress, Hans Selye was also the first to recognize that homeostasis could not by itself ensure stability of body systems under stress. A different tack, focused on cognition, was taken by Lazarus, the eminent and influential Berkeley University psychologist. At a time when psychology tried to understand human behaviour by first understanding simple organisms engaging in simple behaviours learned by associations, rewards, or punishments, Lazarus instead emphasized the importance of studying cognition, which he extended into stress and coping. In parallel with these stress concepts, during the years, neuroendocrine advances revealed the physiological substrate for homeostasis, allostasis, and the stress response mechanisms. The autonomic nervous and the hypothalamic-pituitary-adrenocortical (HPA) systems affect the afferent and efferent limbs of the stress response in vertebrates and are also central to maintaining homeostasis and effecting allostasis (Fink, 2009).
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5.2 Definition and Classification of Stress

The term “stress”, as it is currently used was coined by Selye in 1936, who defined it as “the non-specific response of the body to any demand for change”. Threats to well-being, whether physical or psychological, are components of life experience. Individuals differ markedly, however, in the frequency with which they experience stressful life events and their vulnerability or resilience to stressful challenges (Herman et al., 1995). Stress, although often studied as a psychological construct, may be viewed from a biological perspective (Dantzer, 1991). Accordingly, stress responses are composed of the activation of neurobiological systems that help preserve viability through change or allostasis (McEwen and Seeman, 1999). Although necessary for survival, the effects of frequent physiological stress responses may increase the risk of future physical and mental health problems (Gunnar and Quevedo, 2007).

The Diagnostic and Statistical Manual of Mental Disorders (DSM-V) recognizes two stress disorders: acute stress disorder and posttraumatic stress disorder (PTSD). For the diagnosis of acute stress disorder, the individual, while experiencing the trauma or after the event, must have at least three of several dissociative symptoms, such as a subjective sense of numbing, detachment, or absence of emotional responsiveness; reduction in awareness of surroundings; depersonalization; or dissociative amnesia. Following the trauma, the traumatic event is persistently re-experienced, the individual avoids stimuli that may arouse recollections of the traumatic event, and they have anxiety or increased arousal. PTSD is defined as a condition in which a traumatic event is persistently re-experienced in the form of intrusive recollections, dreams, or dissociative flashback episodes. Cues to the event lead to distress and are avoided, and there are symptoms of increased arousal; the full symptom picture must be present for more than one month, and the disturbance must cause clinically significant distress or impairment in social, occupational, or other areas of functioning (Fink, 2009).

5.3 Neuroanatomy and Physiology of Stress

Stress responses in mammals are affected by two distinct, but interrelated systems: the sympathetic-adrenomedullary (SAM) (Frankenhaeuser et al., 1986) system and the hypothalamic-pituitary-adrenocortical (HPA) (Stratakis and Chrousos, 1995) system. The SAM system is a component of the sympathetic division of the autonomic nervous system, releasing epinephrine (adrenaline) from the medulla or centre of the adrenal gland. Increases in circulating epinephrine facilitate rapid mobilization of metabolic resources and orchestration of the fight/flight response (Jansen et al., 1995). The HPA system, in contrast, produces glucocorticoids (cortisol in humans, corticosterone in rodents; hereafter GCs) which are steroid hormones. Unlike epinephrine, which does not cross the blood-brain barrier to a significant degree, the brain is a major target of GCs (Bohus et al., 1982). Also unlike epinephrine, GCs production takes some time (approximately 25 minutes to peak levels), and many of its impacts on the body and brain occur through the changes in gene expression (de Kloet et al., 1991). Consequently, the impacts of GCs are slower to develop and continue for longer periods (De Kloet et al., 1996). The role of the HPA system in stress is complex, and its functions are not fully captured by reference to the fight/flight response (Sapolsky, 2000). Regulation of both the SAM and HPA systems converges at the
level of the hypothalamus, which integrates autonomic and endocrine functions with behaviour (Palkovits, 1987). Furthermore, inputs to the hypothalamic nuclei that orchestrate HPA and SAM responses to psychosocial stressors involve cortico-limbic pathways (Gray and Bingaman, 1996).

5.4 Stress: social behaviour, and resilience

Social stress, a common stressor readily translated across species, is a recurrent factor in the life of all social species (von Holst, 1998). The effects of stress exposure and consequent trajectory depend on the nature of the stressor, the severity, duration (acute vs. chronic), sex/gender, genetics, timing of exposure (early life, adolescence, adulthood or aging) as well as the perception of the stressor by the individual, for example, stressor controllability dramatically affects resilience versus vulnerability as an outcome (Amat et al., 2010; Lucas et al., 2014; Maier and Watkins, 2005). There are three main categories of social stressors. Life events are defined as abrupt, severe life changes that require an individual to adapt quickly. Chronic stressors are defined as persistent events, which require an individual to make adaptations over an extended period of time. When stress becomes chronic, one experiences emotional, behavioural, and physiological changes that can put one under greater risk for developing a mental disorder and physical illness. Understanding the mechanisms underlying stress-induced disturbances will ultimately allow for improved clinical therapies and possible preventative strategies to decrease the incidence of these disorders.

Resilience refers to a person's ability to adapt successfully to acute stress, trauma or more chronic forms of adversity. A resilient individual has thus been tested by adversity (Rutter, 2006) and continues to demonstrate adaptive psychological and physiological stress responses, or “psychobiological allostasis” (Charney, 2004; McEwen, 2003). Resilience has been linked to being able to perceive stressful events in less threatening ways, promoting adaptive coping strategies; such cognitive reappraisal allows individuals to re-evaluate or reframe adverse experiences in a more positive light (Feder et al., 2009; Southwick et al., 2005).

Although many of the above psychological characteristics cannot be measured in animals, some behavioural traits associated with resilience have been identified. In numerous animal models, rodents display a range of responses to stress: at one extreme are active or “fight-flight” responses (for example, attempts to escape and aggression), and at the other extreme are passive responses (for example, freezing and submission). Active-coping animals are often considered to be resilient, based on numerous functional end points, whereas their more passive counterparts are not; however, both types of responses can be seen as adaptive depending on the particular context (Korte et al., 2005).

In rodents, acute stress typically leads to reduced social behaviours and increased aggression, including antisocial behaviours such as bite counts that exceed species-typical levels (De Almeida et al., 2002; Takahashi et al., 2012). This fits with the concept of acute stress as a 'flight or fight' response and implies that brief acute stressors mobilize resources to cope with the situation (Sandi and Haller, 2015). Chronic stress reduces social motivation and social interactions in a variety of sociability tests (van der Kooij et al., 2014; Wood et al., 2003). However, although chronic stressors generally reduce sociability, social isolation stress actually enhances social interest.
Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide (Van den Berg et al., 1999), probably because long-term deprivation from social contacts increases interest in social partners. Aggressiveness is increased by chronic physical stressors including chronic unpredictable mild stress (Mineur et al., 2003), restraint or immobilization (van der Kooij et al., 2014; Wood et al., 2003; Yohe et al., 2012). By contrast, chronic social stressors that involve fighting that leads to defeat and subordination have been shown to downregulate aggressiveness in various species. Conversely, repeated victories — which are accompanied by reduced physiological stress responses but which can be considered stressful because they involve recurrent exposure to social conflicts — may result in exacerbated and abnormal aggression (Miczek et al., 2013; Nephew and Bridges, 2011).

Stress models that cover a range of neurodevelopmental periods have been applied to investigate the long-term impact of stress on adult social behaviours. Social motivation (sociability) was disrupted in adulthood by prenatal (de Souza et al., 2013), neonatal (Franklin et al., 2011; Yu et al., 2013) and juvenile (Márquez et al., 2013; Naert et al., 2011; Vidal et al., 2011) exposure to stressors. Prenatal stress (de Souza et al., 2013), neonatal stressors -maternal separation (Wei et al., 2013) and early deprivation (Jia et al., 2009) - and peripubertal exposure to physical stressors (Márquez et al., 2013) inhibited social interactions in adulthood. Juvenile social stressors (post-weaning social isolation) (Workman et al., 2011) and early subjugation (Womack et al., 2004) either did not affect this aspect of social behaviour or, in one study (Shimozuru et al., 2008), increased adult social interactions. Early life stressors decrease measures of social motivation, reduce the expression of social behaviours, increase aggressiveness and promote the development of antisocial features, but the specific consequences depend on the timing and type of the early stressor. Although these changes can be problematic for human individuals and societies, from an evolutionary perspective they may be interpreted as mechanisms through which early adversity prepares the organism to endure similar adversities later in life (Gluckman et al., 2007).

5.5 Stress impact on memory function

Stress is a potent modulator of cognitive function in general, and more precisely, of learning and memory processes (de Kloet et al., 1999; McEwen and Sapolsky, 1995; Sandi, 2004). Stress effects are frequently regarded as deleterious to cognitive function; but there are many instances in which neural function and cognition are either facilitated by stress (de Kloet et al., 1999; Joëls et al., 2006), or even not affected (Beylin and Shors, 1998; Warren et al., 1991). This great variability can be explained by the “intensity” of the stressor (Cordero et al., 1998) or internal hormonal reactions (Baldi and Bucherelli, 2005; Conrad, 2005; Joëls, 2006). Another important factor is stress “duration,” with distinct effects frequently induced by single versus repetitive stress or stress hormones activation, and not only at the cognitive level, but also when evaluating brain structure and function (Pecoraro et al., 2005; Pinnock and Herbert, 2001; Sandi and Loscertales, 1999). Also important to empathize is the memory phase at which stress acts: consolidation is generally facilitated and retrieval generally impaired under stress conditions (Roozendaal, 2002, 2003). Psychological factors, notably stressor controllability and predictability that are well known to be key mediators of the psychophysiological impact of stress (Das et al., 2005; Mineka, 1985) have also to be mentioned addressing the variability of stress on memory processes. Convergent
evidence indicates that experiencing uncontrollable stress has deleterious effects on further information processing (Maier and Watkins, 2005). To clarify the outcome of stress in memory function is important to take into account the existence of individual differences, with gender appearing as a strong modulator of such interactions (Bowman et al., 2003; Luine, 2002; Shors, 2004). Certainly relevant to understand how stress affects cognition is the relevance of the context in which stress is experienced, that is, whether stress is or not contingent to the particular information processing under study (de Kloet et al., 1999; Joëls et al., 2006; Sandi, 1998).

Chronic stress was also proposed to compromise the hippocampus, a well-known region in the brain important for memory processing (Eichenbaum, 1997; O'Keefe, 1978). In the hippocampus, chronically activating the stress response can produce maladaptive changes, which have been postulated to contribute to disease (de Kloet et al., 2005; McEwen and Wingfield, 2003; Smith, 1996). A transition into maladaptive changes includes dendritic remodelling resulting in reduces dendritic arbors in CA3 neurons; dendritic retraction has been observed in other brain regions following ten to 21 days of repeated stress. When chronic stress continues for 4 weeks, CA1 and dentate gyrus neurons express dendritic retraction (Sousa et al., 2000). Prefrontal cortical neurons also express dendritic retraction following 1 to 3 weeks of stress (Brown et al., 2005; Radley et al., 2004). Chronic stress-induced CA3 dendritic remodelling has been proposed to be a maladaptive response because it is associated with susceptibility to damage and cognitive dysfunction (McEwen, 2016).

Although the majority of studies on chronic stress have focused on structural changes within the hippocampus, chronic stress has opposite effects in the basolateral nucleus of the amygdala (BLA), where it increases the dendritic complexity of neurons (Vyas et al., 2002; Vyas et al., 2004), suggesting that chronic stress facilitates memory under emotionally arousing situations.

5.6 Chronic Stress Paradigm in rodents

Different animal models have been developed that use chronic stress to induce neuroendocrine and central nervous changes that might reflect also behavioural changes in rodents. The chronic stress paradigms are considered to have a greater etiological relevance and face validity in mimicking depression, anxiety and other diseases.

5.6.1 Chronic Social Defeat Stress

Social stressors have proven to be potent across a wide range of species. Social behaviour is complex and varies with the behavioural test chosen, and whether focal individuals are tested with familiar or novel conspecifics, with same- or opposite-sex individuals, or with familiar or unfamiliar strains. Widely used models of social stress in rodents include social subordination, crowding, isolation, and social instability (Figure 4).

Social rejection is used as a potent experimental stressor (Kirschbaum et al., 1993), because individual's position in the social hierarchy has profound implications for health and well-being (Adler et al., 1994; Sapolsky, 2005). The social defeat test procedure involves the daily exposure to a novel, physically superior aggressor for a defined period of time and results in significantly reduced display of social interaction
Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide and increased anxiety-like behaviour of the defeated animal (van Bokhoven, 2011). Dominance maybe assured by size, prior history of winning, strain of the resident, and/or prior housing differences (Martinez et al., 1998). Social defeat is typically used as a stressor in male rodents, for whom dominance is easier to quantify and aggressive interactions related to home territory are presumed more salient (Beery and Kaufe, 2015). In the short-term, social defeat produces changes in heart rate, hormone secretion, and body temperature, with longer-term impacts on a wide variety of additional outcomes including activity, social behaviour, drug preference, disease susceptibility and others (Martinez et al., 1998; Peters et al., 2012; Sgoifo et al., 1999). Social defeat stress has profound effects on hippocampal morphology and function (Buvvalda et al., 2005; McEwen, 2012; McEwen and Magarinos, 2001; Mirescu and Gould, 2006). These effects include reduction in hippocampal volume (Czéh et al., 2001) related to dendritic remodelling and reduced neurogenesis (Gould et al., 1998; Magariños et al., 1996). After prolonged exposure to social defeat stress (and stressful conditions in general), animals often show an altered brain structure and cognition (McEwen, 1999; Sandi and Pinelo-Nava, 2007), that is said to increase the risk of developing neuropsychiatric disorders (de Kloet et al., 2005; Nemeroﬀ et al., 2006)). At the molecular level, BDNF has been reported to play an important role in social defeat stress (Berton and Nestler, 2006). As the key downstream regulatory factor, CREB is a constitutively expressed regulatory nuclear transcription factor involved in not only stress, but also individual development and synaptic plasticity (Guzowski and McGaugh, 1997; Sakamoto et al., 2011).

Social defeat is considered to be one of the most robust animal models of stress-induced mood-related illnesses (Berton and Nestler, 2006). Compared to other animal models, the social defeat paradigm has higher face, predictive, and ethological validity, which results in enduring behavioural and neurobiological changes that mimic several symptoms of the human condition (Iñiguez et al., 2014).

5.6.2 Crowding and Isolation

Housing density affects rodent behaviour, and both crowded and isolated social environments have been used as stressors in rodents. Crowding is a naturalistic stressor especially for social or gregarious species that relates to high population density and resource competition in the field. In house mice, several studies have shown that crowding can impair reproductive function and may be part of population size regulation (Christian, 1971; CHRISTIAN and LEMUNYAN, 1958). Increased group size is associated with greater dispersal consistent with a “social competition” hypothesis (Quirici et al., 2011). Social crowding has been shown to affect many different physiological outcomes in rodents. These include changes in organ weights, hormone secretion, HPA reactivity, pain sensitivity, telomere length, and cardiac outcomes (Gadek-Michalska, 2003; Gamallo et al., 1986; Grippo et al., 2011; Kotrshal et al., 2007; Puzserova et al., 2013; Tramullas et al., 2012). At the opposite extreme, solitary housing can be a potent stressor for social species. Solitary housing produces an “isolation syndrome” particularly in females, consisting of hyperadrenocorticism, reduced body weight, altered blood composition, and enhanced pain responsiveness among other outcomes (Hatch et al., 1965; Valzelli, 1973). These changes coincide with alterations in behaviour including aggression, mating behaviour, learning, and pain sensitivity (Valzelli, 1973).
5.6.3 Social Instability

Some studies employ both crowding and isolation in alternation as a model for chronic social instability (Haller et al., 1999; Herzog et al., 2009). In the social instability stress paradigm, uncontrollability is modelled by alternating isolation and crowding phases and by the rotation of animals among social groups during the crowding phase (Herzog et al., 2009). Social instability has particularly been used as a social stressor for female rats, for whom crowding and social defeat are not always effective stressors (Palanza et al., 2001). In the crowding phase, different social groups consisting of different numbers of males and females are formed. Females exposed to this variable social environment show increased adrenal weight, increased corticosterone secretion, decreased thymus weight, and reduced weight gain relative to females housed in stable male e female pairs (Haller et al., 1999). Social instability also induced dysregulation of the hypothalamic-pituitary-gonadal (HPG) axis and reduced sucrose preference and food intake (Herzog et al., 2009).

5.6.4 Chronic Restrain Stress

Chronic restrain is induced by placing rodents into a well ventilated transparent tube for a few hours per day for 2-3 weeks. The animals display depressive-like behaviour (Haenisch et al., 2009) and also impairment of hippocampal neurogenesis, down-regulation of neurotrophins, such as BDNF, and alteration in synaptic plasticity markers (Pollak et al., 2010; Veena et al., 2009). However, daily exposure to the same kind of stressor at the same time is experienced as predictable mild stress, which may actually improve mood and enhance hippocampal neurogenesis in mice (Parihar et al., 2011) so that the relevance of this paradigm is questionable if not associated with other stressors.

5.7 Neuronal histamine: an insight on stress

Histamine release is a sensitive indicator of stress (Taylor and Snyder, 1971; Verdière et al., 1977), and chronic restraint and/or metabolic stress are among the most potent activators of histamine neurons in the TMN (Miklós and Kovács, 2003). Distinct subgroups (E4-E5) of hypothalamic histamine neurons respond to immobility, foot shock, hypoglycemia, and dehydration, suggesting a functional heterogeneity of histaminergic TMN neurons (Miklós and Kovács, 2003). TMN neurons are influenced by a number of neuroendocrine signals (Gotoh et al., 2005) and may integrate exteroceptive and interoceptive state cues in the control of stress-induced arousal. Histamine mediates the stress-induced neuroendocrine hormone surges of ACTH, β-endorphin, and AVP from the pituitary (Kjaer et al., 1992) and controls stress-related activity of aminergic systems, including serotonin-, norepinephrine-, dopamine-, and acetylcholine-containing neurons. As an integral part of the neural networks generating autonomic patterns histamine neurons interfere with AVP- and CRH-positive sympathetic command neurons (Krout et al., 2003) in the PVN and LHA (Whitcup et al., 2004) to influence sympatho-adrenal outflow, cardiovascular functions, and complex stress-related behaviours such as 'flight or fight' or grooming. Histamine injections in the PVN activate the HPA axis through CRH release. Moreover, both hista-
Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide mine and CRH are released from mast cells in the leptomeninges and along brain capillaries during systemic stress emphasizing the intricate interaction between histamine and CRH, and the nervous and immune system (Esposito et al., 2002).
Chapter 6
Histamine and the gut-brain axis

The bidirectional signalling between the gastrointestinal tract and the brain is vital for maintaining homeostasis and is regulated at the neural (both central and enteric nervous systems), hormonal and immunological levels. Perturbation of these systems results in alterations in the stress-response and overall behaviour (Rhee et al., 2009). The high co-morbidity between stress-related psychiatric symptoms such as anxiety with gastrointestinal disorders including irritable bowel disorder (IBS) and inflammatory bowel disorder (IBD) (Cámara et al., 2009; Mawdsley and Rampton, 2006) are further evidence of the importance of this axis. However, increasing evidence also suggests that the enteric microbiome greatly impacts on gut-brain communication (Cryan and O'Mahony, 2011). The necessary communication processes are based on neurotransmitters, neuropeptides, cytokines, hormones, growth factors (among others), which mediate the relationship between the immune system and the CNS (Downing and Miyan, 2000). Also, luminal probiotic bacteria may alter behaviour and brain biochemistry in a variety of ways even in the absence of changes in the inflammatory status of the host (Bercik, 2011). Bacterial products could enter the circulation to pass the blood-brain barrier if they are sufficiently small and lipophilic, or they might enter the brain at the circumventricular organs where the barrier is diminished. Since prior vagotomy abolishes behavioural and brain biochemical changes induced by certain probiotic bacteria (Bercik et al., 2011; Bravo et al., 2011), afferent vagal signalling is a necessary condition for the central effects of these neuroactive microorganisms.

Under stress conditions, the brain may influence the composition of the gut microbiota (Bailey and Coe, 1999) via the hypothalamus–pituitary–adrenal (HPA) axis, which regulates cortisol secretion, affecting immune cells activity both locally in the gut and systemically. Stress can affect such equilibrium (Glaser and Kiecolt-Glaser, 2005), leading to allergic reactions, inflammatory responses and predisposition to infection. In turn, gut microbiota, and probiotic agents can alter the levels of circulating cytokines, which in turn can have a marked effect on several brain functions (Duerkop et al., 2009; Forsythe and Bienenstock, 2010).

Recent advances in understanding the molecular and systems biology of the gut-brain axis have suggested novel candidates such as gut lipid sensors, and have revealed novel roles for lipid sensing in the control of nutrient availability (Schwartz, 2011) and in cognitive functions. One of these sensors is the endocannabinoid oleoylethanolamide.
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6.1 Oleoylethanolamide

The ability to monitor the nutrient composition of food allows animals to generate neural and endocrine feedback signals that adapt behaviour and metabolism to environmental fluctuations in food availability. Being able to sense dietary fat is especially important, from an adaptive perspective, because of the high energy density of this nutrient and the essential role of lipids as building blocks of cell membranes and precursors for hormones and other bioactive molecules. The need to secure appropriate amounts of fat and optimize its storage and utilization provides a plausible evolutionary basis for the existence of multiple lipid-sensing mechanisms throughout the mammalian body (Piomelli, 2013). New evidence suggests an important role for a different class of signalling molecules – the amides of fatty acids (FAs) with ethanolamine (fatty acid ethanolamides, FAEs). The FAEs were first isolated from plant and animal tissues more than 50 years ago (Bachur et al., 1965; Kuehl, 1957) and were found to possess strong anti-inflammatory activity (Lambert et al., 2002). The discovery of anandamide (AEA), a polyunsaturated FAE led to uncovering the function served by FAE family molecules as regulators of food intake (Rodríguez de Fonseca et al., 2001). This family includes the monounsaturated analogue oleoylethanolamide (OEA) (Piomelli, 2013), which, although sharing similar biosynthetic pathways (Okamoto et al., 2007) with AEA, exerts opposite effects on feeding regulation and lipid metabolism (Romano et al., 2014).

A calcium-dependent N-acyltransferase activity (NAT) catalyses the transfer of a fatty acyl group from the sn-1 position of phosphatidylcholine to the amine group of phosphatidylethanolamine, producing a family of N-acylphosphatidylethanolamine (NAPE) species. NAPEs containing oleic acid at the amine position generate OEA upon hydrolysis of their distal phosphodiester bond. This reaction is catalysed by a NAPE-selective phospholipase D (NAPE-PLD), which produces phosphatidic acid (PA) as a by-product (Figure 6). OEA is hydrolysed into oleic acid and ethanolamine by either of two structurally unrelated enzymes: FA amide hydrolase (FAAH) or N-acylthanolamine acid amidase (NAAA) (Piomelli, 2013).

In particular, OEA is synthesized in the upper part of the small intestine, upon the absorption of lipids from the diet (Fu et al., 2007; Li et al., 2015). Endogenous OEA itself is nutritionally regulated specifically by intestinal lipid administration. Food deprivation reduces OEA biosynthesis in the small intestine, while intestinal lipid infusions, but not equicaloric carbohydrate or protein infusions, increase proximal small intestinal OEA levels (Schwartz et al., 2008). Interestingly, the biosynthetic pathway responsible for lipid stimulated endogenous OEA production also generates intra-intestinal oleic acid, bringing full circle an extrinsic food-derived lipid signal to a de novo intra-intestinal oleate signal (Schwartz et al., 2008). Feeding suppressive effects of small intestinal lipid infusions are mediated in part by a local regulatory network within upper small intestine: nutritional lipid is translocated into the luminal tissue and stimulates gut OEA and oleic acid biosynthesis (Schwartz, 2011).

Peripheral OEA administration reduces food intake by increasing the latency to feed and by prolonging the interval between two successive meals (Fu et al., 2005; Gaetani et al., 2003; Rodríguez de Fonseca et al., 2001). OEA hypophagic actions cannot be attributed to stress or malaise because OEA does not produce behaviours that are indicative of fear or anxiety, does not change plasma corticosterone levels, and does not induce conditioned taste aversion in rats (Fu et al., 2005; Proulx et al.,
The hypogastic actions of OEA depend on the feeding state of the animal. In free-feeding rats, OEA decreases meal frequency without altering meal size; by contrast, the compound simultaneously reduces these two parameters in food-deprived animals (Gaetani et al., 2003).

The effects of OEA on food intake appear to be mediated in part by Peroxisome Proliferator Activated Receptor (PPAR)-alpha, a nuclear receptor that is also implicated in the mechanism that allows liver cells to gauge the levels of circulating FAs (Pawar and Jump, 2003). OEA fails to suppress feeding in PPAR-alpha deficient mice, while PPAR-alpha agonists mimic the feeding inhibitory effects of exogenous OEA administration (Rodríguez de Fonseca et al., 2001). Gut vagal afferents also contribute to the ability of OEA to reduce feeding, as vagal capsaicin application blocks OEA induced satiety (Rodríguez de Fonseca et al., 2001). Consistent with this finding, OEA also rapidly depolarizes capsaicin-sensitive cell bodies of vagal afferent neurons in the nodose ganglion. This activation is likely mediated by the transient receptor potential vanilloid type 1 (TRPV1) receptor, as it is blocked by TRPV1 antagonists and is absent in TRPV1 null mice (Wang et al., 2005). OEA also acts as an agonist for GPR119 (Suardiaz et al., 2007), a G protein-coupled receptor that recognizes a broad panel of lipid molecules in addition to OEA (Hansen et al., 2012). GPR119 is expressed in intestinal endocrine L-cells, which secrete glucagon-like peptide-1 (GLP-1), and intraluminal infusions of OEA were found to increase circulating GLP-1 levels in rats (Lauffer et al., 2009). Nevertheless, the observation that genetic deletion of TRPV1 or GPR119 in mice does not alter the anorexic effects of OEA strongly argues against a direct involvement of these receptors in OEA-induced satiety (Lan et al., 2009; Lo Verme et al., 2005).

Intraperitoneal injections of OEA stimulate transcription of c-Fos in the Nucleus of the Solitary Tract (NST) and in peptide-secreting neurons of the paraventricular (PVN) and supraoptic (SON) nuclei of the hypothalamus (Gaetani et al., 2010; Rodríguez de Fonseca et al., 2001) and increase the expression of the neuropeptide oxytocin, but not vasopressin. Moreover, pharmacological blockade of central oxytocin receptors abrogates the hypogastic effects of OEA, implying that release of oxytocin in the hypothalamus and/or other regions of the brain may be a key effector of OEA-induced satiety (Gaetani et al., 2010).

OEA recruits other neurotransmitter systems in the brain to reduce food intake. Recently in our laboratory it was shown that histamine deficiency significantly attenuates OEA-induced hypophagia (Provensi et al., 2014). As previously reported in this thesis, brain histamine affects feeding behaviour, it is fundamental for appetitive and aversive responses during motivated behaviour, and blockade of histamine H1R in the hypothalamus is believed to be responsible for the weight gain and metabolic dysregulation associated with the clinical use of atypical antipsychotics (Kim et al., 2007). Provensi and co-workers (2014) demonstrated that lack of central histamine dampens OEA-induced increase of c-Fos expression in oxytocin PVN neurons. Therefore, OEA requires the integrity of the brain histamine system to fully exert its hypogastic effect and the finding establish new functional connections between peripherally acting hypogastic signals and brain histamine neurotransmission (Provensi et al., 2014).

OEA exert also other functions mediated at least in part by the intestine-brain connections. By using the inhibitory avoidance and the Morris water maze test, Campolongo et al. (2009) found that i.p. administration of OEA after behavioural training
Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide strongly improved the retention of these tasks. This effect was blocked by infusions of the local anaesthetic lidocaine into the NST or the β-adrenergic antagonist propranolol into the BLA, indicating that the signal launched by peripheral OEA gains access to the CNS and strengthens memory consolidation by stimulating noradrenergic activity in the BLA (Campolongo et al., 2009). This suggest that OEA signalling initiates an integrated response in which satiety induced by a fat-rich meal coincides temporarily with enhanced encoding of information about the spatial and emotional context in which the meal was consumed highlighting the importance of metabolic peripherally acting signals in the regulation of higher brain function (Berthoud and Morrison, 2008).

Other behavioural effects of OEA include antidyskinetic effect as assessed by using a hemiparkinsonian model of PD in mice with 6-OHDA striatal lesion. OEA treatment reduced axial, forelimb and orolingual dyskinetic symptoms, as well as contralateral rotations induced by 6-OHDA (González-Aparicio and Moratalla, 2014). Also, OEA reduced spontaneous locomotor activity and attenuated psychomotor activation induced by cocaine, an effect that does not seem to be mediated by PPARα receptor (Bilbao et al., 2013). OEA’s central effects were also tested in depressive-like behaviour by using two animal models: the chronic unpredictable mild stress (CUMS) and tail suspension/forced swim tests. OEA treatment normalized sucrose preferences, rearing frequencies, prefrontal cortex and hippocampal atrophy and reversed the abnormalities of BDNF and MDA levels and SOD activities in the hippocampus and prefrontal cortex, as well as changes in serum levels of ACTH, CORT, and T-AOC in CUMS, demonstrating antioxidant properties and normalization of the hyperactivity in the hypothalamic–pituitary–adrenal axis (HPA) (Jin et al., 2015). In the TST/FST, OEA treatment decreased the immobility time demonstrating an antidepressant-like effect and also and increased cerebral levels of NE and 5-HT regulating central monoamine neurotransmitters (Yu et al., 2015a).
Aim of the study

The main scope of my study is to understand the role of the histaminergic system in the homeostatic and behavioural effects of oleoylethanolamide (OEA), a fat sensing hormone normally secreted by enterocytes in the jejunum. Our working hypothesis is that brain histamine serves as a relay station that elaborates peripheral and central signals to allocate the relevance and adequate arousal to perform the appropriate behaviour.

For the purpose of clarity, the manuscript is divided in three parts concerning different aspects of the interaction of the histaminergic system and oleoylethanolamide.

In the first part of the work we brain histamine is necessary for OEA to exert its effects on emotional memory. The results appear in the publication Provensi et al., 2017 (Provensi et al., 2017) that I co-authored.

In the second part, I investigated whether histamine has a role in OEA’s antidepressant-like effects. In this regard, our research group recently demonstrated that selective serotonin reuptake inhibitors require the integrity of the brain histamine system to exert their preclinical responses (Munari et al., 2015). Consequently we hypothesised a possible relationship between antidepressant-like effect of OEA (Jin et al., 2015) and brain histamine. To this end, we evaluated the repeated and chronic i.p. administration of OEA in mice unable to synthesize HA due to disruption of the histidine decarboxylase gene (HDC-KO) or to injection of alpha-fluoromethylhistidine compared to wild type (WT), and saline i.c.v. injected controls in the tail suspension test. The phosphorylation level of cyclic AMP-response element binding protein (pCREB), a major player in the molecular mechanisms of antidepressant treatment was also evaluated by Western Blot analysis in prefrontal cortex and hippocampus, two regions critically involved in depression and antidepressant effects of drugs.

In the third part of the thesis we investigated the effect of behavioural effects of OEA in chronic social defeat stress (CSDS)-induced cognitive impairments. In 1971 it was reported that brain histamine turnover was altered by stress conditions (Taylor and Snyder, 1971). Stress affects a constellation of physiological systems in the body and evokes a rapid shift in many neurobehavioral processes. In our hypothesis histamine may detect stress-induced signals from the periphery. As the influence of OEA on stress reactivity is not currently addressed, we evaluated the effect of OEA in HDC-KO and wild type mice subjected to the CSDS. In CSDS paradigm experimental mice are exposed to social and physical conflict by aggressive member of the same species resulting in both physical and emotional stress for 21 days, while control non-stressed mice will be handled daily. The behavioural repertoire of the animals has been assessed at the end of the CDSC, using a battery of tests that are comprehensive of several domains affected by stress: social behaviour, mood, anxiety and cognition.
1.1 Materials and Methods

1.1.1 Animals and Drugs

Male Wistar rats (3 months old, 300–330 g) purchased from Envigo (Bresso, Italy) were housed in the animal facility of Ce.S.A.L (Università di Firenze) in a temperature-controlled room (22±1°C) with a 12-h light/dark cycle (light on 7:00 AM to 7:00 PM), at a constant temperature and humidity with standard diet (4RF21; Mucedola s.r.l., Milan, Italy) and freely available water. All procedures were conducted in accordance with the Council Directive of the European Community (2010/63/EU) of the Decreto Legislativo Italiano 26 (13/03/2014) and National Institutes of Health guidelines on animal care and were approved by veterinarian supervision.

Alpha-FMHis was synthesized at Johnson & Johnson Laboratories (kind gift of Dr. Nicholas Carruthers), pyrilamine was purchased from Sigma-Aldrich (UK), and zolantadine and OEA from Tocris Bioscience (UK). OEA was dissolved in saline/polyethylene glycol/Tween80 (90/5/5, v/v), whereas zolantidine and pyrilamine were dissolved in saline. All other reagents and solvents were of HPLC grade or the highest grade available (Sigma).

1.1.2 Surgery

One week after arrival, rats were anaesthetized (75 mg/kg ketamine plus 10 mg/kg xylazine) and placed on a stereotaxic frame (Stellar; Stoeling Co., Wood Dale, IL). A stainless-steel cannula (22 gauge) was implanted in the lateral ventricle and fixed to the skull by using dental cement according to the following coordinates (Paxinos et al., 1998) in mm: AP=−0.9; L=−1.5; DV=−2.6 and used for α-FMHis/saline administration. Rats were also implanted bilaterally with 22-gauge guide cannulae 1 mm above the BLA according to the following coordinates from bregma (Paxinos et al., 1998) in mm: AP=−2.8; L=±4.9; DV=+7.6. Animals were allowed 7 days to recover from surgery before behavioural procedures and were handled once daily before the experimental day.
Results
Part I
Histaminergic Neurotransmission as a Gateway for the Cognitive Effect of Oleoylethanolamide in Contextual Fear Conditioning

1.1 Materials and Methods

1.1.1 Animals and Drugs

Male Wistar rats (3 months old, 300–330 g) purchased from Envigo (Bresso, Italy) were housed in the animal facility of Ce.S.A.L (Università di Firenze) in a temperature-controlled room (22±1°C) with a 12-h-light/dark cycle (light on 7:00 AM to 7:00 PM), at a constant temperature and humidity with standard diet (4RF21; Mucedola s.r.l., Milan, Italy) and freely available water. All procedures were conducted in accordance with the Council Directive of the European Community (2010/63/EU) of the Decreto Legislativo Italiano 26 (13/03/2014) and National Institutes of Health guidelines on animal care and were approved by veterinarian supervision.

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1.1.3 Infusion Procedure and Experimental Groups

At the time of drug microinfusions, the animals were gently restrained by hand, and the injection needle (30 gauge) was fitted tightly into the guides, extending 1 mm from the tip of the guide cannulae. The injection needle was connected to a 10-μL Hamilton microsyringe, and the infusions were performed at a rate of 0.5 μL/30 s. The infusion cannula was left in place for an additional 60 seconds to minimize backflow. The entire bilateral infusion procedure took approximately 90 seconds. Alpha-FMHis (5 mM, 1 µL) was infused i.c.v. 24 hours before contextual fear training, and controls received equal volumes of sterile saline. Zolantidine (0.1 μM, 0.5 μL side) or pyrila-mine (0.9 μM, 0.5 μL side) were infused intra-BLA bilaterally immediately after training. OEA (10 mg/kg) was injected i.p. 10 minutes after fear conditioning, while controls received equivalent volumes of vehicle.

1.1.4 Contextual Fear Conditioning

Contextual fear conditioning was induced in a Skinner box module (29 × 31 × 26 cm, Modular Operant Cage; Coulbourn Instruments Inc.), equipped with a grid floor connected to a shock-delivery apparatus (Modular Operant Cage/Grid Floor Shocker E13-08; Coulbourn Instruments) and placed in an acoustically insulated room at 20±1°C. The number of the electric shocks and the inter-shock interval duration was predetermined by a stimulus programming device (Scatola di comando Arco 2340, Italy). Illumination inside the room was 60 lux. The rat was left undisturbed for 3 minutes and subsequently six, 1-seconds 0.8-mA electric footshocks were adminis-tered at 30-second intervals. The footshock intensity was chosen according to previous published data from our laboratory (Benetti et al., 2013). This is a strong enough footshock to guarantee retention at 72 hours postacquisition without inducing general-ization (Baldi et al., 2004). The rat was removed 2 minutes after the end of the stim-ulation, therefore spending a total time of 8 minutes inside the conditioning apparatus.

1.1.5 Freezing Measurement

Seventy-two hours after conditioning, rats were again placed inside the condition-ing apparatus in the soundproof room and left undisturbed for 6 minutes. The rats’ behaviour was recorded by means of a closed-circuit television system by an experimenter unaware of the animal’s treatment. Freezing was defined as the complete ab-sence of somatic motility, with the exception of respiratory movements. Measure-ments were performed with a stopwatch by personnel unaware of the experimental group each animal belonged to. Total cumulated freezing time (i.e., total seconds spent freezing during each 6-minute period) was calculated and results expressed in seconds of freezing time. All behavioral tests were performed between 10:00 AM and 12:00 PM to avoid interference with the circadian rhythm (Kamin, 1957).

1.1.6 Histology

The placement of infusion cannulae was verified postmortem. Rats were over-dosed with chloral hydrate and the brains removed and stored in 10% formalin for 10 days. Forty-μm sections were sliced on a cryostat, mounted on gelatine-coated slides,
and then stained with cresyl violet for light microscopic observation. Data from rats in which the cannulae were not correctly positioned were discarded (<5%).

1.1.7 Statistical Analysis

All values are expressed as means ± SEM, and the number of rats used in each experiment is also indicated. The presence of significant treatment effects was determined by a Student’s t test or a 1-way ANOVA followed by Newmann Keuls’ MCT test, as appropriate. For all statistical tests, P<.05 was considered significant.

1.2 Results

1.2.1 Oleoylethanolamide administration increases freezing time of rats submitted to contextual fear conditioning.

In a first set of experiments we evaluated the effect of OEA (10 mg/kg i.p. a dose that does not change motility in the open field, nor anxiety-like responses, (Campolongo et al., 2009) administered within 10 min of contextual fear conditioning in satiated rats. Controls received an equivalent volume of vehicle. Retention test was carried out 72 hrs after training. As shown in figure 7, an unpaired Student’ t test showed that rats given OEA displayed a significant increase of time spent freezing compared with vehicle treated animals (p<0.01).

1.2.2 Histaminergic neurotransmission is required for OEA-freezing enhancements

To evaluate the role of the central histaminergic system in the cognitive effect of OEA, we infused the HDC inhibitor α-FMHis i.c.v. 24 hours prior to fear conditioning. Our research group previously showed that administration of α-FMHis quickly suppressed baseline and histamine H3 receptor antagonist-evoked release of histamine from the TMN of freely moving rats (Benetti et al., 2015), as 180 minutes after injection, histamine release values decreased below the sensitivity of the method. OEA (10 mg/kg i.p.) was injected within 10 minutes after fear conditioning. Controls received saline i.c.v. and vehicle i.p. One-way ANOVA revealed a statistical difference across experimental groups (F3,45=6.756; P<.001; Figure 8). Post-hoc analysis with Newman-Keuls MCT showed that OEA increased the freezing time at retention test with respect to vehicle-treated rats receiving i.c.v. infusion of saline (P<.05). However, the OEA-elicited potentiation of freezing was abolished in brain histamine-depleted animals (P<.01). Hence, the treatment with α-FMHis 24 hours prior to the test prevented the effect of OEA, indicating that the integrity of the central histaminergic system is necessary for the effects of OEA on memory consolidation. The freezing time of rats given vehicle i.p. and of those receiving α-FMHis or saline i.c.v. did not differ significantly at retention, thus indicating that all animals formed a memory trace of the training experience.
Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide

1.2.3 Antagonism of histamine H_1 and H_2 receptors prevents OEA-induced freezing enhancement

Figure 9 shows the effect of blocking H_1 or H_2 receptors in the BLA with local, bilateral infusions (0.5 μL) of the selective H_1 antagonists pyrilamine (0.9 μM) or the H_2 antagonist zolantidine (0.1 μM), immediately prior to the administration of OEA. Controls received intra-BLA, comparable volumes of saline and i.p injections of vehicle. One-way ANOVA showed significant differences among experimental groups ($F_{5,64} = 8.436; P<.0001$). Rats that received OEA i.p. and saline intra-BLA froze for a significantly longer time at retention compared with saline-/vehicle-treated rats ($P<.05$) in a comparable manner to the control groups in Figure 8. However, when either H_1 or H_2 receptors were blocked, OEA administration did not increase the freezing time ($P<.001$). Freezing time of rats that received vehicle i.p. and pyrilamine or zolantidine in the BLA was not significantly different from the freezing time of saline-/vehicle-treated rats.

1.3 Summary of Results (Part I)

Emotional arousal enhances consolidation of memory traces, a homeostatic response of our organism that is modulated by stress hormones (McGaugh and Roozendaal, 2002). There is extensive evidence derived from observations in both experimental animals and humans that the amygdala is indispensable to enable the acquisition and retention of lasting memories of emotional experiences. The compelling evidence led McGaugh and his collaborators (McGaugh, 2004) to propose that the activation of the BLA is fundamental in the establishment of an arousal state triggered by fear as unconsciously occurs in humans, (LeDoux, 2014) and that arousal is a major component in the establishment of post-training memory consolidation.

By activating peripheral PPAR-α OEA engages the vagus nerve and activates the NTS that in turn provides the amygdala, together with the locus coeruleus (LC), with a dense supply of norepinephrine, and modulates long term memories (Campolongo et al., 2009).

The present results show that OEA increases memory expression when tested in the contextual fear conditioning paradigm, by increase the freezing time of treated animals compared to vehicle injected rats. Depletion of histamine in the brain with infusion of α-FMHIs that blocks HDC, or intra-BLA infusions of antagonists of the H_1 or H_2 receptor prevent the freezing-enhancing effects of OEA. Therefore, the histaminergic system in the amygdala exerts a permissive role on the memory-enhancing effects of OEA. When histaminergic neurotransmission is compromised by α-FMHIs-induced depletion OEA effects on memory expression is prevented. It is relatively surprising that both H_1 and H_2 receptor antagonists produce the same results. The arrangement of these receptors on BLA neurons is not known, and explanations may only be speculative. Presumably, when strong aversive stimuli are used, all histaminergic inputs to the BLA need to be blocked to prevent freezing-enhancing compounds such as OEA to exert their effect.

The concept of several neurotransmitter systems contributing to emotional memory consolidation within the same brain region is indisputable (Izquierdo et al., 1992; Izquierdo et al., 2016; McGaugh, 2004). We think that the emotional arousal that is generally considered indispensable for good memory consolidation of fear tasks

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is provided by both the noradrenergic and histaminergic transmission in the BLA. In our hypothesis histamine released in the amygdala gates the effects of noradrenaline on BLA neurons, hence blocking either neurotransmitter within the BLA would lead to similar behavioural outcomes. In this case, H1 and H2 receptors work synergistically presumably activating common intracellular pathways and their blockade leads to similar behavioural outcomes.

Taking into consideration the results presented in this thesis and previous evidence (Campolongo et al., 2009), OEA appears to increase the expression of BLA-dependent emotional memory regardless of the type of behavioural response and adverse situation. From a translation point of view, few human studies published so far showed that blood concentrations of N-acylethanolamides like OEA and other endocannabinoids increase in response to acute stress in healthy human volunteers (Dlugos et al., 2012), and subjects affected by Post-traumatic stress disorder (PTSD) have significantly higher plasma concentrations of OEA (Hauer et al., 2013; Schaefer et al., 2014). These results are based on small and dis-homogeneous cohort of patients, nonetheless they may have diagnostic relevance. In this regard, the results presented in this thesis, suggest that activation of the histaminergic system in the BLA has a “permissive” role on the memory enhancing effects of OEA. In particular, targeting the H1 or H2 receptor with classical clinically approved antihistamine compounds may modify the expression of emotional memory and reduce dysfunctional aversive memories as found in phobias and PTSD.
Part II

Oleoyl ethanolamide induces antidepressant-related responses by targeting PPAR-α and recruiting the histaminergic neurotransmission.

2.1 Materials and Methods

2.1.1 Animals and Drugs

Male CD1 mice (25-35 g; Harlan, Italy), histidine decarboxylase knockout (HDC-KO) mice and wild type (WT) littermates (background sv129), grown in the Centro Sabulazione Animali di Laboratorio (CeSAL), Università di Firenze were used for behavioural and biochemical experiments. Genetically modified animals were routinely genotyped using the PCR protocol previously described (Provensi et al., 2014). Peroxisome proliferator activated receptor-alpha knockout (PPAR-α-KO) mice and wild type (WT) littermates (background C57Bl/6), were used for behavioural and biochemical experiments. Wild-type and PPAR-α-KO (B6.129S4-SvJae-Pparatm1Gonz) mice previously backcrossed to C57BL6 mice for 10 generations were bred in Centro Stabulazione Animali di Laboratorio, Università Federico II di Napoli, and the colony was established and maintained by heterozygous crossing. Mice were genotyped as described on the supplier webpage (http://jaxmice.jax.org). Animals were housed in humidity and temperature-controlled room (22-24 °C), allowed free access to food (4RF21; Mucedola s.r.l., Milan, Italy) and water, and kept on a 12-h light/dark cycle (lights start at 7:00 AM). All the experiments were conducted between 9:00 AM and 4:00 PM. OEA and vehicle were dissolved as previously reported (Result part I). Imipramine (IMI) was dissolved in saline. Acute neuronal histamine deprivation in CD1 mouse was induced by injection intracerebroventricular (i.c.v) of 5µL of α-fluoro-methyl-histamine solution (1µg/µL). The control group received an injection of 5 µL of saline ic.v..

2.1.2 Behavioral Experiments

2.1.2.1 Tail Suspension Test (TST)

Mice were fixed with a 1 cm piece of adhesive tape from the tip of the tail, suspended 50 cm from the ground, at least 30 cm away from the objects that may surround them and the behavior was filmed by a camera positioned in front of the apparatus for 6 min. The immobility time was manually timed by a researcher unaware of the treatment. OEA was administered following two different regimens: 10 mg/kg
Part II
Oleoylethanolamide induces antidepressant-related responses by targeting PPAR-α and recruiting the histaminergic neurotransmission

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Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide (OEA) repeated administration, 3 i.p. injections in 24 hours and 10 mg/kg i.p. OEA subchronic with 1 daily i.p. injection for 8 days before challenging the mice in TST 1 h after the last treatment. The test was carried out in a soundproof room with low light and was recorded using a video camera (Digital Cam 34x, Samsung). The time that mice remained immobile (immobility time) in the last 4 minutes of test was analyzed off-line by an operator unaware of the pharmacological treatments.

### 2.1.2.2 Open Field test

One hour after the last treatment, animals were positioned in a corner of an openfield arena (60cm×70cm) and the general motor activity was assessed in a 5 minutes session. The covered distance (cm) was evaluated using Smart 2.5 software.

### 2.1.3 Neurochemical experiments

#### 2.1.3.1 Western Blot analysis

Male HDC-KO, PPAR-α-KO and WT mice received i.p. injections of Vehicle, OEA (10 mg/kg) or imipramine (IMI) (10 mg/kg) daily for 8 days and 1 h later they were challenged in the TST, immediately after the test, they were sacrificed. The brains were dissected on ice, and the cortices and hippocampi isolated immediately. The cortices and the pooled hippocampi (left and right) were individually homogenized in 200 μL ice-cold lysis buffer containing protease and phosphatase inhibitors (50mM TrisHCl pH 7.5, 50mM NaCl, 10 mM EGTA, 5mM EDTA, 2mM NaPP, 4 mM PNFF, 1 mM Na3VO4, 1.1 mM PMSF, 20μg/μL Leupeptin, 50 μg/μL Aprotinin, 0.1% SDS) and centrifuged at 12000 rpm at 4°C for 15 minutes. The supernatant was collected, and total protein levels were quantified using the Pierce BCA Protein Assay (Thermo Scientific, USA). Homogenates were diluted in a mix of lysis buffer and loading buffer 2x (50mM Tris pH 6.8, 100mM DTT, 10% Glycerol, 1% Bromophenol Blue and 2% SDS) and boiled for 10 minutes. Aliquots containing 40 μg total proteins were resolved by electrophoresis on a 10% SDS-polyacrylamide gel (SDS-PAGE) and transferred onto polyvinylidene difluoride (PVDF) membranes (Immobilon Transfer Membranes, Millipore, USA). Blots were blocked in Tris-buffered saline, pH 7.6 containing 0.1% of Tween 20 (TBS-T) and 5% non-fat dry milk (Bio-Rad Laboratories, USA) for 2 h at room temperature and then incubated overnight at 4°C with monoclonal antibodies against α-phospho-CREB (Ser 133) or αCREB (both from Cell Signaling Technology, USA) diluted 1:1000 in TBS-T containing 5% bovine serum albumin or 5% non-fat dry milk, respectively. Immunodetection was performed with secondary antibodies (anti-rabbit IgG conjugated to horseradish peroxidase, Cell Signaling Technology, USA) diluted 1:5000 in TBS-T containing 1% of non-fat dry milk. Membranes were washed in TBS-T and then reactive bands were detected using enhanced chemiluminescence (Luminata Crescendo, Millipore, USA). Quantitative densitometric analysis was performed using the QuantityOne analysis software (Bio-Rad). For each sample the ratio of αpSer133-CREB/αCREB densities was calculated and then all the individual rates were expressed as a percentage of the average of ratios obtained from control groups.
2.1.4 Statistical Analysis

All values are expressed as means ± SEM, and the number of mice used in each experiment is also indicated. The presence of significant treatment effects was determined by using a 1-way ANOVA followed by Newmann Keul’s MCT test, as appropriate. The level of significance was set at P ≤ 0.05.

2.2 Results

2.2.1 Oleoylethanolamide systemic administration exerts antidepressant-like effect by recruiting histaminergic neurotransmission

We assessed the possible antidepressant-like effect of repeated or sub-chronic OEA treatment in the TST response of HDC-KO mice and their wild type littermates (WT). Imipramine was used a positive control.

Figure 10 shows the effect of different doses of OEA administered 3 times in 24 hrs. OEA increased immobility time in WT mice at both doses used, whereas it was ineffective in HDC-KO mice. A 2-way ANOVA revealed an overall significant difference between groups F(genotype x treatment) 3,69 = 5.026, p<0.005; F(treatment) 3,69 = 12.85, p<0.0001; F(genotype) 1,69 = 4.466 p<0.05. Bonferroni’s post hoc test showed that OEA treatment at both doses significantly decreased the immobility time of WT mice exposed to the TST (5 mg/kg, p < 0.01; 10 mg/kg, p<0.001), whereas no significant effects were observed in HDC-KO mice. Imipramine decreased the immobility time of both genotypes (WT and HDC-KO, p<0.001) (Figure 10A).

When tested after sub-chronic treatments ANOVA revealed an overall significant difference between HDC-KO and WT mice. Either OEA, imipramine or vehicle were administered i.p. for 8 consecutive days, and the last injection 1 hr before the test a 2-way ANOVA showed significant differences between groups (F(genotype x treatment) 3,57 = 3.362, p<0.05); F(treatment) 3,57 = 7.418, p<0.001; F(genotype) 1,57 = 10.34 p<0.01). Bonferroni’s post hoc analysis revealed that OEA treatment at all doses tested significantly reduced the immobility time of WT mice (5 mg/kg and 10 mg/kg OEA, p<0.001), whereas no effect on immobility time was observed in HDC-KO mice. Imipramine reduced the immobility time of both genotypes (p<0.001) (Figure 10B).

To avoid biases related to compensatory mechanisms due to chronic depletion of histamine, we assessed the effects of OEA in CD1 mice that received i.c.v. infusion of the HDC suicide inhibitor α-FMHIs. In analogy to the results observed in chronically histamine deprived mice, also in α-FMHIs-treated mice, ANOVA revealed an overall significant difference between groups (F(7,92) = 14.35; P<0.0001); In analogy to HDC-KO mice, OEA did not modify the immobility time of α-FMHIs treated mice (97.4 ± 25.0 s) compared to vehicle-treated mice (149.1 ± 18.9 s; Figure 12), whereas imipramine remained effective regardless of the presence of brain histamine (F(icv x treatment) 3,79 = 4.707, p<0.01); F(treatment) 3,79 = 43.62, p<0.0001; F(icv) 1,79 = 5.007, p<0.05). Significant differences for OEA treatments by Bonferroni’s post hoc test were p<0.001 (Figure 10C).
Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide

2.2.2 OEA-induced increase in cortical and hippocampal CREB phosphorylation is reduced in HDC-KO mice

We compared the effect of OEA (10 mg/kg), imipramine (10 mg/kg) and vehicle-treated HDC+/+ and HDC−/− mice. Two-way ANOVA showed an overall significant difference between groups (F(genotype x treatment) 2,21 = 4.14, p<0.05; F(treatment) 2,21 = 14.05, p=0.0001; F(genotype)1,21 = 11.56, p<0.05). Bonferroni’s post hoc test found significant differences between groups (p<0.001; Figure 11). OEA increased significantly pCREB in the hippocampus of HDC+/+ mice compared with vehicle treated animals (Bonferroni’s post hoc test p<0.01; Figure 11A). However, OEA treatment was ineffective in HDC−/− mice (Figure 11A), as pCREB levels were not different from those of vehicle treated mice. Similar to the effect in the hippocampus, OEA increased pCREB levels in the frontal cortex of HDC+/+ mice (Figure 11B), but not in the cortex of HDC−/− mice (2-way ANOVA, F(genotype x treatment) 2,24 = 3.78, p<0.05; F(treatment) 2,24 = 6.78, p<0.01; F(genotype)1,24 = 3.57, p=0.07). Imipramine administered at 10 mg/kg for 8 days increased pCREB in the hippocampus and cortex of both genotypes with respect to controls (p<0.05).

2.2.3 Oleoylethanolamide systemic administration reduced immobility time by targeting PPAR-α

To explore if PPAR-α are required for the antidepressant-like effect of OEA, PPAR-α-KO mice and wild type littermates were tested in the TST after repeated or sub-chronic OEA (10 mg/kg) treatments as shown in Figure 12. A 2-way ANOVA was conducted to examine the effect of genotype and repeated (3 x in 24hrs) treatment on immobility time that revealed an overall significant difference between groups (F(genotype x treatment)1,29 = 5.924, P<0.05; F(treatment)1,29 = 11.80, p<0.01; F(genotype)1.29 = 0.8, p=0.3). Bonferroni post-hoc test showed that OEA treatment significantly decreased the immobility time of PPAR-α+/+ mice (p<0.001), but not of PPAR-α−/− mice (Figure 12A).

When tested after OEA sub-chronic treatments (daily injection for 8 days) ANOVA revealed an overall significant difference between groups (F(genotype x treatment)1,27 = 9.147; P<0.01; F(treatment)1,27 = 2.94, p=0.09; F(genotype)1.27 = 0.8, p=0.28). Post hoc analysis revealed a reduction in immobility time in WT mice treated with 10 mg/kg OEA compared with VEH treated animals (p<0.01; Figure 12B). In PPAR-α-KO male mice no differences were observed between 10 mg/kg OEA and VEH treated animals (Figure 12B).

2.2.4 OEA-induced increase in cortical and hippocampal CREB phosphorylation is reduced in PPAR-alpha KO mice

Figure 13 shows CREB phosphorylation at Ser 133 residue in cortical and hippocampal homogenates of normal and PPAR-α deficient mice treated sub-chronically with OEA (10mg/kg) or vehicle. Two-way ANOVA revealed an overall difference in hippocampal (F(genotype x treatment)1,30 = 4.943, p<0.05, F(treatment)1,30 = 10.61, p<0.01; F(genotype)1.30 = 0.6, p=0.44) and cortical (F(genotype x treatment)1,26 = 3.032, F(treatment)1,26 = 3.514, p=0.09; F(genotype)1.26 = 0.02, p=0.8) homogenates between WT and PPARα-KO mice. Post-hoc comparisons showed a significant increase in pSer133CREB/CREB...
ratios after OEA treatment when compared with vehicle-treated control animals in both the hippocampus (Bonferroni’s post hoc test p<0.01; Figure 13A) and cortex (Bonferroni’s post hoc test p<0.05; Figure 13B). No significant changes in CREB phosphorylation were observed in PPAR-α KO mice that received OEA or vehicle.

2.2.5 General motility

To exclude possible effects of the various treatments and genotypes on spontaneous locomotor activity that may have affect the immobility time in the TST, mice were exposed to the open field and motor activity recorded for 5 min. As shown in figure 14, no differences were observed between experimental groups.

2.3 Summary of the Results (Part II)

The present results indicated that repeated treatments with OEA reduced the immobility time in the tail suspension test when compared with vehicle treated HDC+/+ mice, but this effect is not observed in HDC−/− mice. All antidepressants effectively reduce immobility in this test (Lucki, 2001). Imipramine, a classic tricyclic antidepressant, reduced the immobility time in both genotypes. To understand if brain histamine is necessary for the behavioural effects of OEA, and to exclude compensatory mechanisms of HDC−/− mice, we also evaluated the behaviour in mice pharmacologically deprived of histamine by using an i.c.v injection of α-FMH, a suicide inhibitor of HDC. Similarly to what we observed in HDC−/− mice, a significant reduction in the immobility time was observed in WT mice treated with OEA, but not in α-FMH-treated mice. OEA treatments did not modify the exploratory behaviour in the open field test performed by each experimental group indicating that the effect observed in the TST is not related to alterations in the motor activity.

We tested the hypothesis that different levels of CREB phosphorylation - a transcription factor that serves as a convergence point for multiple classes of antidepressant drugs (Carlezon et al., 2005) - would explain the differences between genotypes in the observed behavioural tests. Our results also indicate that histamine signalling is necessary for OEA to trigger CREB phosphorylation. By using a sub-chronic treatment regimen, we observed a good correlation between behavioural and biochemical data. OEA and imipramine treatment reduced the immobility time and increased CREB phosphorylation in cortical and hippocampal homogenates of HDC+/+ animals, whereas in HDC−/− mice OEA did not change immobility time, nor CREB phosphorylation levels. Furthermore, imipramine increased CREB phosphorylation and reduced immobility time independently of genotype. Our results mirror the observation of Munari et al., (2015) who found that selective serotonin reuptake inhibitors (SSRI) such as citalopram reduced the immobility time and increased pCREB of HDC+/+ mice, but was ineffective HDC−/− mice even though their serotonergic system is functional and the pCREB pathway is not compromised (Munari et al., 2015). Thus, our biochemical results indicate that the lack of CREB phosphorylation may contribute to OEA inefficacy in histamine-deficient mice.

Where histamine and OEA interact to exert their behavioural and neurochemical effects remains to be established. In this thesis, I provide evidence that OEA targets the PPAR-α to exerts its effects on the TST and CREB. To this end we evaluated the behaviour of PPAR-α+/+ and PPAR-α−/− mice. We observed that repeated treatments
Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide with OEA reduced the immobility time and increased pCREB in WT mice, but not in PPAR-\(\alpha^-\) mice.

In conclusion, our results confirm that OEA may improve animal depression-like behaviours as first observed by Jin et al., (2015) (Jin et al., 2015), but we unravelled a novel mechanism by which OEA achieves its effect, namely by recruiting the brain histaminergic system. We believe that activation of the histaminergic system, at least in the hippocampus and prefrontal cortex, is necessary for OEA-induced CREB phosphorylation and consequently the behavioural output.

Although OEA shows potential as an antidepressant that reverses cerebral functional abnormalities and ameliorates depression, further research should be performed on the specific pathways by which OEA affects the histaminergic system and behavioural responses.
Part III
Histaminergic involvement in Oleoylethanolamide protection on the cognitive decline induced by chronic social stress in mice

3.1 Materials and Methods

3.1.1 Animals and Drugs

Male CD1 mice (9 weeks of age; Charles River, Italy), histidine decarboxylase knockout (HDC-KO) mice and wild type (WT) littermates (background c57Bl/6), grown in the Centro Sabulazione Animali di Laboratorio (CeSAL), Università di Firenze were used for behavioural experiments. Genetically modified animals were routinely genotyped using the PCR protocol previously described (Provensi et al., 2014). Animals were housed in humidity and temperature-controlled room (22 - 24 °C), allowed free access to food (4RF21; Mucedola s.r.l., Milan, Italy) and water, and kept on a 12-h light/dark cycle (lights start at 7:00 AM). All the experiments were conducted between 9:00 AM and 4:00 PM. OEA and vehicle were dissolved as previously reported (Result part I). The OEA or Vehicle treatments started at ten days before the end of stress procedure.

3.1.2 Chronic Social Defeat Stress Paradigm

Prior to the social defeat stress, we selected the aggressive CD1 resident. CD1 mice were screened for aggressive behavior: latency of first attack was monitored and dominance status of mice was visually determined. Mice were deemed dominant if they displayed aggressive behavior toward their opponent (another CD1) such as tail rattling, chasing, biting and fight-attacks. Mice were submissive if they displayed defending and avoidance behaviour such as escaping, defensive response, upright posture and defensive immobility. Mice with latencies to attack of >30 s were not selected. The selected mice were those that were the most aggressive, most dominant, the heaviest and those which had the lowest latency to attack, for two consecutive days.

Chronic Social Defeat Stress (CSDS), was a 21-days social defeat adapted from previous studies (Bartolomucci et al., 2001; Keeney et al., 2006). In this paradigm the C57Bl/6 mice were daily introduced in the cage of CD1 aggressive resident until the first attack. A 3-min cut-off for latency of attack was imposed in order to maintain the interaction between mice short. Thereafter, mice were physically separated by a perforated transparent divider and remained in sensory (olfactory and visual) contact for 2 h. After that the divider is removed and the second attack occur. Stresses mice were
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single housed during the entire period of stress. Control mice were left undisturbed housed in their own home cage with other control mouse. This chronic stress was combined with overcrowding conditions because housing density affects rodent behaviour (Beery and Kaufer, 2015) and crowded social environments have been used as stressors in the protocol. Food consumption and body weight were evaluated daily during the protocol.

3.1.3 Social Interaction Test

The social interaction test is used to assess social behaviour (Berton and Nestler, 2006). This is a two-step procedure. In the first 2.5 min session, the experimental C57BL/6 mouse is allowed to freely explore an open field arena (41 cm length × 32 cm width × 15 cm height). Along one side of the arena is a rectangular (9.5 cm length x 7.5 cm width x 5 cm height) wire cage that remains empty during the first trial (target absent condition). The C57BL/6 mouse is then removed from the testing arena for 1 minute (into a home cage), and a novel CD1 male mouse is placed into the wire cage. In the second 2.5 min trial (target present condition), the experimental C57BL/6 mouse is reintroduced into this arena now containing a social target (unfamiliar CD1 mouse) within the rectangular wire cage. Time (sec) spent in the interaction zone (surrounding wire cage) in the presence of the social target (Iñiguez et al., 2014), served as dependent variables.

3.1.4 Novel object recognition test

Object recognition paradigm measures a form of memory based on short and un-repeated experiments without any reinforcement, such as food or electric shocks (Ennaceur and Delacour, 1988). Object recognition is a one-trial task, and does not involve the learning of any rule, being entirely based on the spontaneous exploratory behavior of rodents toward objects. Mice were placed in a white polyvinylchloride box (70 × 60 cm and 30 cm high) with a grid floor that is easily cleaned and illuminated by a 75-W lamp suspended 50 cm above the box. The objects to be discriminated were grey polyvinyl shapes: cubes of 8 cm side, pyramids and cylinders of 8 cm height. The object recognition task consisted of a training phase (T1) and a testing phase (T2). Twenty-four h prior to T1, they were habituated for two 10 min-session to the experimental apparatus in the absence of any object. Each mouse was subjected to the procedure separately and care was taken to remove any olfactory/taste cues by cleaning carefully the arena and test objects between trials. On the day of the experiment, the mouse was placed for 5 min into the test arena facing the same direction and in the same position in the presence of two identical plastic objects (cubes) (T1). The behaviour of mice was videotaped, and the time spent actively exploring the objects was measured. Exploration was defined as sniffing or touching the objects with the nose and/or forepaws. Sitting on or turning around the objects was not considered exploratory behaviour. T2 was performed 1 h after T1, during which, each mouse was again placed in the test arena for 5 min in the presence of one of the familiar objects and a novel object. The position of the objects (left/right) was randomized to prevent bias from order or place preference. Mice were placed in their home cage between trials. The behaviour of mice during T2 was videotaped, and the exploration periods of the familiar (F) and the new object (N) were measured. Care was taken to avoid
place preference and olfactory stimuli by randomly changing the role (familiar or new object), and the position of the two objects during T2, and cleaning them carefully. Object recognition was carried out in an insulated room to avoid any noise that could impair the performance of the mouse.

### 3.1.5 Statistical analysis

All values are expressed as means ± SEM, and the number of mice used in each experiment is also indicated. The presence of significant treatment effects was determined by a 2-way ANOVA followed by Bonferroni MCT test, as appropriate. The level of significance was set at \( P \leq 0.05 \). Statistical analysis was performed using GraphPad Software. The data of the object recognition test are expressed as a percentage of time spent exploring each object during T1 and T2.

### 3.2 Results (Part III)

#### 3.2.1 Chronic Social Defeat Stress induced body weight gain and increased food consumption in OEA or VEH treated mice compared to controls

Figure 15 shows the effect of OEA (10 mg/kg) or vehicle i.p. injections in mice submitted to the chronic social defeat stress (CSDS) on body weight and food consumption. Two-way ANOVA revealed no statistical significance between groups in WT mice (Figure 15A) nor in the HDC-KO mice (Figure 15B). Figure 15C shows the cumulative food consumption of WT mice treated with OEA or VEH and subjected to the CSDS protocol. Two-way ANOVA revealed no statistical significance between groups. Figure 15D shows the cumulative food consumption of HDC-KO mice treated with OEA or VEH and subjected to the CSDS protocol. Two-way ANOVA revealed an overall significant difference between groups (\( F_{\text{treatments}}^{(2,336)} = 198.9, \ P < 0.0001 ; \ F_{\text{days}}^{(20,336)} = 952.76, \ P < 0.0001 \)).

#### 3.2.2 Oleoylethanolamide reduces social avoidance induced by social defeat stress

Figure 16 shows the effect of OEA systemic administration to stressed animals submitted to CSDS compared to controls, in the social avoidance test. OEA was administered for the 10 days preceding the test. One-way ANOVA revealed an overall significant difference between groups (\( F_{2,17}^{(2,17)} = 20.36 \ # p < 0.05 \) OEA vs VEH; **p < 0.001 OEA vs NON-STRESSED; ***p > 0.0001 VEH vs NON-STRESSED). Stressed mice spent less time in the interaction with the CD1 mouse compared to non-stressed control mice, therefore, time spent interacting was influenced by stress exposure. OEA treatment increased the interaction time when compared to controls treated with vehicle (Figure 16A). On the contrary, ANOVA revealed no significant differences among HDC-KO regardless of the treatment either (OEA or vehicle). However, an overall significant difference was revealed between stressed and non-stressed groups (1-way ANOVA and Bonferroni post-hoc test; \( F_{2,18}^{(2,18)} = 10.23 \ * p < 0.05 \ **p < 0.001 \); Figure 16B).
Histaminergic neurotransmission as a gateway for the effects of Oleoylethanolamide

3.2.3 Oleoylethanolamide improves the performance in the object recognition test of WT stressed mice, but not of HDC-KO mice

Figure 17 shows the effect of OEA administration on mice performances in the object recognition test. No significant group effects were detected during T1 (two-way ANOVA and Bonferroni post-hoc test) (Figure 17A). During T2, given 1 hour after the T1, Two-way ANOVA revealed an overall difference between groups (F (objects x treatment)2,38=1.915 P=0.16; F (objects)1,38=18.10, p<0.0001; F (treatment)2,38=6.330e-014, p=1). Non-stressed control mice spent more time exploring the familiar object (p<0.001). Similarly, stressed mice treated with OEA spent significantly more time exploring the new object than the familiar one (p<0.01). On the contrary, two-way ANOVA revealed no significant differences in the exploration of the two objects of stressed mice treated with vehicle (Figure 17B). In HDC-KO mice no significant group effects were detected during T1 (two-way ANOVA and Bonferroni post-hoc test) (Figure 17C). During T2 Two-way ANOVA revealed an overall statistical difference between groups (F (objects x treatment)2,31=4.536, P<0.01; F (objects)1,31=0.14, p=0.7; F (treatment)2,31=0.005, p=0.99). Non-stressed HDC-KO mice spent more time in the exploration of the novel object compared to familiar one (p<0.05). No differences in the exploration of the familiar and the novel object were observed in stressed HDC-KO mice treated with OEA or Vehicle (two-way ANOVA and Bonferroni post-hoc test) (Figure 17D).

3.2.4 CSDS did not affect motility of mice tested in the open field

When evaluated in the open field test ANOVA revealed no significant differences between experimental groups after treatments with OEA or VEH in stressed WT mice compared to controls (Figure 18). Similarly, HDC-KO mice motility was not affected by stress or treatments.

3.3 Summary of the Results (Part III)

Stress and traumatic events are increasingly recognized as risk factors for mental disorders in particular for depression, anxiety disorders, and post-traumatic stress disorder (Musazzi et al., 2017; Selten and Cantor-Graae, 2005, 2007; Selten et al., 2013) (PTSD). Several rodent models of neuropsychiatric disorders use chronic stress, measuring behavioural and neurochemical readouts at the end of the protocols.

The present data indicate that the social defeat stress induce social avoidance behaviour as observed by reduction of interaction time in the social interaction test between stressed and non stressed mice. It has been widely reported that defeated animals of different species spend on average significantly less time in close proximity to a social target in a social interaction test as compared to non-defeated animals (Berton and Nestler, 2006; Dadomo et al., 2011; Hollis et al., 2010). The treatment with OEA reduce the social avoidance in WT compared to vehicle treated mice and non stressed controls. To understand if brain histamine is necessary for the behavioural effects of OEA, we also evaluated the behaviour in HDC-KO mice. The OEA effect in reducing social avoidance was not observed in histamine deprived animals.

Stress is a biologically significant factor that, by altering brain cell properties, can disturb cognitive processes such as learning and memory (Kim and Diamond,
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Conclusions and Discussion

The histamine system is a pleiotropic system. Histamine plays a major role in the regulation of autonomic functions, including energy balance, sleep and regulation of body temperature (Haas and Sergeeva, 2012) and is crucial in controlling arousal and cognition (Köhler et al., 2011).

HA-synthesizing neurons are strongly conserved throughout vertebrate phylogeny, from zebrafish to humans reflecting the important roles they play in many aspects of physiology and behavior (Haas and Panula, 2003; Haas et al., 2008; Panula and Nuutinen, 2013; Passani et al., 2007; Schwartz et al., 1991). In mammals, histaminergic neurons are only found in the tuberomamillary nucleus (TMN) of the posterior hypothalamus (Panula et al., 1984; Watanabe et al., 1984) and extend a widespread and diffuse, network of unmyelinated fibers throughout the central nervous system, including interconnections with other arousal-related neuromodulatory systems (Panula and Nuutinen, 2013).

Disruptions of histaminergic neurotransmission have been implicated in a variety of neuropsychiatric disorders (Baronio et al., 2014; Shan et al., 2015a; Shan et al., 2015b). The histamine system has been suggested as a possible target for the treatment of psychiatric disorders, and drugs that modulate this system have been especially proposed as cognitive enhancers (Passani and Blandina, 2011; Tiligada et al., 2011). Therefore, a better understanding of the histamine system would be of great use for the development of new, much needed, pharmacological treatments for psychiatric disorders.

Oleoylethanolamide is an endogenous lipid mediator that inhibits feeding and in rodents, intestinal OEA levels increase about threefold upon refeeding, a response that may contribute to the induction of between-meal satiety (Piomelli, 2013). OEA was also identified in the gastrointestinal tract of Python molurus (Astarita et al., 2006) suggesting that this lipid messenger may be widespread among vertebrate groups and may represent an evolutionarily ancient means of regulating energy intake.

It was recently demonstrated in our laboratory that brain histamine mediates the central effects of a signal molecule produced in the intestine, namely oleoylethanolamide (OEA). OEA is released by the enterocytes in response to high fat intake and reduces eating (Fu et al., 2005) by indirectly activating a subpopulation of histaminergic neurons (Provensi et al., 2014).

In my thesis, I explored other potential interactions between these two phylogenetically ancient systems in the pursuit of unexplored neuronal mechanisms that may shed light on the mode of action of psychoactive agents and possibly lead to the development of new drugs.
Conclusions and Discussion

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OEA was shown to affect not only feeding behaviour, but also the consolidation of aversive memories (Campolongo et al., 2009). Here, I demonstrated that the activation of the histaminergic system in the amygdala of rats is necessary for OEA to modulate the consolidation of fear memories (Provensi et al., 2017).

As reported in the present thesis we also demonstrated that OEA has a potential antidepressant effect since it reduces immobility time in the TST, a paradigm widely used to predict antidepressant properties of several compounds. The reduction of immobility time occurred in normal but not in histamine-deprived mice. In a parallel fashion, OEA, as other antidepressant compounds elicited CREB phosphorylation in normal mice but not in brain histamine-deprived mice, nor in PPAR-α−/− mice.

Hence, we provided evidence that the molecular target for the antidepressant-like effect of OEA is PPAR-α. The most conservative hypothesis is that activation of peripheral PPAR-α in turn recruit the histaminergic system in a yet unexplored fashion. Understanding the mechanisms of OEA’s antidepressant-like action could improve the understanding of depression and the treatment of a disease that is not completely understood and do not respond adequately to available agents.

The several roles of OEA in brain disorders led us to test its effects also in stress-related disorders. Stress, as previously described, is known to induce physical, behavioural and neuropathological outcomes. Here we showed that the maladaptive behavioural responses to chronic stress, such as memory impairments and a depression-like phenotype may be prevented by OEA sub-chronic treatment. Once again, we evaluated the role histaminergic system play in such effect, considering its involvement in stress responses (Ito, 2000). Here, I reported that the beneficial effect of OEA to prevent stress-induced behavioural impairments are only achieved in mice with an intact brain histamine system. Therefore, in this paradigm as well, brain histamine is necessary for OEA’s neurological effect.

Our current working hypothesis is based on the concept that histaminergic neurons act as a relay station that integrate peripheral signals, organize them into a coherent output to selected brain regions hence sending information to other brain areas in order to create a neuro-circuitry responsible for central OEA’s effects.

OEA could be just an example, but many other peripheral signals may influence brain functions using this mechanism. This hypothesis implies that histamine neurons in the TMN are organized in distinct subpopulations differently regulated and innervating specific brain areas to allow multiple processes.

The identification of signalling pathways connecting the periphery with the central nervous system is fascinating and holds great promise for the development of new therapeutics, as targeting peripheral systems may provide a more easily accessible therapeutic strategy for pharmaceutical treatment.
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PPAR

of OEA is PPAR

Hence, we provided evidence that the molecular target for the antidepressant

end of the histamine system is the H3 receptor. Therefore, we decided to focus our

studies on these receptors. In this chapter, we report our findings on the role of histamine

H3 receptors in the regulation of stress and depression-related behavior.

The several roles of OEA in brain disorders led us to test its effects also in stress

We first investigated the effect of OEA on stress-induced behavioral changes using the

forced swim test (FST). This paradigm is widely used to assess the antidepressant activity

of experimental compounds. Our results showed that OEA significantly reduced the immobility

time occurred in normal but not in histamine

backed mice. In a parallel experi-

ence, OEA, as other antidepressant compounds elicited CREB phosphorylation in

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of aversive memories

OEA could be just an example, but many other peripheral signals may influence

Our current working hypothesis is based on the concept that

that OEA directly activates histaminergic neurons in the hypothalamus and this

response is mediated by the PPARα receptor. In line with this hypothesis, we found that OEA

increased the expression of PPARα in the hypothalamus of mice. These results suggest that

the PPARα receptor is involved in the mechanism of action of OEA.

As reported in the present thesis we also demonstrated that OEA has a potential

long-term effects on stress and depression-related behavior. In particular, OEA was shown to

hibit the consolidation of long-term memories in the Morris water maze task. This

indicates that OEA may be a potential therapeutic agent for the treatment of depression

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Figures

Figure 1. The histaminergic system in human brain. The histaminergic fibers emanating from the tuberomamillary nucleus project to and arborize in the whole central nervous system (Haas et al., 2008).

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Figure 1. The histaminergic system in human brain. The histaminergic fibers emanating from the tuberomammillary nucleus project to and arborize in the whole central nervous system (Haas et al., 2008).
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Figure 2. Key brain areas involved in the regulation of feeding and their innervation by histaminergic fibres (Panula and Nuutinen, 2013).
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Figure 3. cAMP response element-binding protein/brain-derived neurotrophic factor signal transduction pathway (Wang et al., 2013).

Figure 4. Schematic representation of the levels at which the social environment impacts and reflects the individual (Beery and Kaufer, 2015).
Figure 5. Microbiota–gut–brain (MGB) axis. Direct and indirect pathways support the bidirectional interactions between the gut microbiota and the central nervous system (CNS); involving endocrine, immune and neural pathways (Montiel-Castro et al., 2013).
Figure 5. Microbiota – gut – brain (MGB) axis. Direct and indirect pathways support the bidirectional interactions between the gut microbiota and the central nervous system (CNS); involving endocrine, immune and neural pathways (Montiel-Castro et al., 2013).

Figure 6. Biochemical pathway responsible for the production of OEA (Fu et al., 2011).
Figure 7. OEA (10 mg/kg) or vehicle (VEH) were injected i.p. 10 minutes after contextual fear conditioning. Fear retention was evaluated 72 hours after conditioning. Bars represent mean values ± sem of 8 to 10 rats/group; **P < .01; unpaired t test.
Figure 8. α-FMHiS or saline was infused i.c.v. 24 hours prior to fear conditioning. OEA (10 mg/kg) or vehicle (VEH) were injected i.p. 10 minutes after contextual fear conditioning. Fear retention was evaluated 72 hours after conditioning. Latencies of α-FMHiS groups did not significantly differ from controls. Data are expressed as means ± SEM of 10 to 14 animals for each group; ANOVA and Newman-Keuls posthoc test, *P<.05 vs saline (SAL) controls. ##P<.01 vs OEA/SAL.
Figure 9. Rats received bilateral intra-BLA infusions of zolantidine (ZOL), pyrilamine (PYR), or saline, and OEA or vehicle i.p. immediately after training. Data are expressed as means ± SEM of 9 to 14 animals for each group; ANOVA and Newman-Keuls’ posthoc test, **P < .01 vs SAL/VEH controls; ###P < .001 vs SAL/OEA.
Figure 10. OEA induce antidepressant-like effects in normal but not in histamine-deficient mice in the Tail Suspension Test. (A) Effect of systemic administration of OEA (repeated regimen) or vehicle to WT and HDC-KO male mice (n=5-12). (B) Effect of systemic administration of OEA (10 mg/kg i.p. sub-chronic regimen) or vehicle to WT and HDC-KO male mice (n=5-10). (C) Effect of systemic administration of OEA in saline or α-FMH (5 µg i.c.v.) injected CD1 mice (n=5-12).
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Figure 11. OEA chronic administration increased Ser133CREB phosphorylation in hippocampal (A) and cortical (B) homogenates from normal (WT) but not from histamine-deprived mice (HDC-KO) challenged in the tail suspension test. (n=5; *p<0.05 **p<0.01 ANOVA and Bonferroni post hoc test).

Figure 12. OEA induces antidepressant-like effects in normal but not in PPAR-α-KO mice in the Tail Suspension Test. (A) Effect of systemic administration of OEA (repeated regimen) or vehicle to WT and PPAR-α-KO male mice (n=5-12). (B) Effect of systemic administration of OEA (10 mg/kg i.p. sub-chronic regimen) or vehicle to WT and PPAR-α-KO male mice (n=5-10). (**p<0.01 ***p<0.001 ANOVA and Bonferroni MCT).
Figure 13. OEA chronic administration increased Ser133CREB phosphorylation in and hippocampal (A) cortical (B) homogenates from normal (WT) but not from PPAR-α-deprived mice (PPAR-α KO) challenged in the tail suspension test. (n=7-9; *p<0,05 **p<0,01 Two-way ANOVA and Bonferroni post hoc test).
Figure 14. OEA systemic administration did not change general motor activity in normal, histamine deficient- and PPAR\(\alpha\)-KO mice in the open field test. (A) Effect of systemic administration of OEA (10 mg/kg i.p. repeated regimen) or vehicle to WT and HDC-KO male mice (n=5-11). (B) Effect of systemic administration of OEA (5 or 10 mg/kg i.p. sub-chronic regimen) or vehicle to WT and HDC-KO male mice (n=5-11). (C) Effect of systemic administration of OEA (10 mg/kg i.p.) in saline or \(\alpha\)-FMH (5 \(\mu\)g i.c.v.) treated CD1 mice (n=8-11). (D) Effect of systemic administration of OEA (10 mg/kg i.p. sub-chronic regimen) or vehicle to WT and PPAR-\(\alpha\) KO male mice (n=5-11). (ANOVA and Newman-Keuls post hoc test).
Figure 14. OEA systemic administration did not change general motor activity in normal, histamine deficient and PPAR α-KO mice in the open field test. (A) Effect of systemic administration of OEA (10 mg/kg i.p. repeated regimen) or vehicle to WT and HDC-KO male mice (n=5-11). (B) Effect of systemic administration of OEA (5 or 10 mg/kg i.p. sub-chronic regimen) or vehicle to WT and HDC-KO male mice (n=5-11). (C) Effect of systemic administration of OEA (10 mg/kg i.p.) in saline or α-FMH (5 µg i.c.v.) treated CD1 mice (n=8-11). (D) Effect of systemic administration of OEA (10 mg/kg i.p. sub-chronic regimen) or vehicle to WT and PPAR α-KO male mice (n=5-11). (ANOVA and Newman-Keuls post hoc test).

Figure 15. Effects of the chronic social defeat stress on body weight and food consumption. CSDS (A) induced no significant increase in body weight in OEA and VEH treated stressed mice compared with non-stressed control group (n=3-9). (B) No differences were detected in body weight between experimental group in HDC-KO mice (n=6-7). (C) CSDS induced no significant increase in cumulative food consumption in OEA and VEH treated stressed mice compared with non-stressed control group (n=3-9). (D) CSDS induced a significant increase in cumulative food consumption in OEA and VEH treated stressed HDC-KO mice compared to non stressed controls (n=6-7 **p<0.01 ***p<0.0001).
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Figure 16. Social defeat stress induced social avoidance in WT and HDC-KO mice. (A) Defeated mice spent less time in the interaction compared to controls (**p<0.001 OEA vs NON STRESSED; ***p>0.0001 VEH vs NON STRESSED), OEA treatment increased interaction time in defeated animals compared to VEH treated controls (#p<0.05 OEA vs VEH). (B) HDC-KO defeated mice spent less time in the interaction compared to controls (*p<0.05 **p<0.001). OEA treatment did not increase interaction time in HDC-KO compared to controls.
Figure 16. Social defeat stress induced social avoidance in WT and HDC-KO mice. (A) Defeated mice spent less time in the interaction compared to controls (\( p<0.001 \) OEA vs NON STRESSED; \( *p>0.0001 \) VEH vs NON STRESSED), OEA treatment increased interaction time in defeated animals compared to VEH treated controls (\( #p<0.05 \) OEA vs VEH). (B) HDC-KO defeated mice spent less time in the interaction compared to controls (\( *P<0.05 \) **p<0.001). OEA treatment did not increase interaction time in HDC-KO compared to controls.

Figure 17. Effect of OEA on mice performances in the object recognition test. (A) Time spent in the exploration of Object A and B during T1 (One-way ANOVA and Bonferroni’s MCT). (B) T2 was performed 1 h after training. Results are calculated as individual percentage of time spent exploring familiar (black columns) and novel (white columns) objects. One-way ANOVA and Bonferroni’s MCT revealed an overall significant difference in the exploration of novel object compared to familiar one in non stressed controls and OEA treated stressed mice (means ± S.E.M. of 3-9 animals per experimental group. ****\( p<0.0001 \); **\( p<0.01 \), vs. respective familiar object). (C) Time spent in the exploration of Object A and B during T1 in HDC-KO mice (One-way ANOVA and Bonferroni’s MCT). (D) Percentage of time spent exploring familiar (black columns) and novel (white columns) objects in stressed and non stressed control mice. One-way ANOVA revealed an overall significant difference in the exploration of novel object compared to familiar one in non stressed controls (±SEM of 6-7 animals per experimental group. *\( p<0.05 \)).
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Figure 18. CSDS did not change general motor activity in normal and histamine deficient mice in the open field test. (A) Effect of CSDS in WT animals treated with OEA (10 mg/kg i.p.) or vehicle compared to non stressed controls (n=3-7). (B) Effect of CSDS in HDC-KO animals treated with OEA (10 mg/kg i.p.) or vehicle compared to non stressed controls (n=6-7) (ANOVA and Bonferroni post hoc test).
Figure 18. CSDS did not change general motor activity in normal and histamine deficient mice in the open field test. (A) Effect of CSDS in WT animals treated with OEA (10 mg/kg i.p.) or vehicle compared to non stressed controls (n=3-7). (B) Effect of CSDS in HDC-KO animals treated with OEA (10 mg/kg i.p.) or vehicle compared to non stressed controls (n=6-7) (ANOVA and Bonferroni post hoc test).
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