Role of neuroinflammation in the emotional and cognitive alterations displayed by animal models of obesity

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Obesity is associated with a high prevalence of mood disorders and cognitive dysfunctions in addition to being a significant risk factor for important health complications such as cardiovascular diseases and type 2 diabetes. Identifying the pathophysiological mechanisms underlying these health issues is a major public health challenge. Based on recent findings, from studies conducted on animal models of obesity, it has been proposed that inflammatory processes may participate in both the peripheral and brain disorders associated with the obesity condition including the development of emotional and cognitive alterations. This is supported by the fact that obesity is characterized by peripheral low-grade inflammation, originating from increased adipose tissue mass and/or dysbiosis (changes in gut microbiota environment), both of which contribute to increased susceptibility to immune-mediated diseases. In this review, we provide converging evidence showing that obesity is associated with exacerbated neuroinflammation leading to dysfunction in vulnerable brain regions associated with mood regulation, learning, and memory such as the hippocampus. These findings give new insights to the pathophysiological mechanisms contributing to the development of brain disorders in the context of obesity and provide valuable data for introducing new therapeutic strategies for the treatment of neuropsychiatric complications often reported in obese patients.

Keywords: obesity, anxiety, depression, memory impairments, inflammation, neuroinflammation, indoleamine 2,3-dioxygenase, GTP-cyclohydrolase 1

Introduction

The prevalence of obesity has been steadily and alarmingly increasing worldwide for decades, fostering a rise in serious obesity-related outcomes, particularly cardiovascular diseases and metabolic disorders that contribute to a significant rise in mortality. In addition, obesity is also increasingly linked to a number of psychopathologies including mood disorders and cognitive dysfunctions (Luppino et al., 2010; Francis and Stevenson, 2013). The incidence of depressive symptoms (up to 30%) is much higher in obese subjects than in normal weight age-matched population (Roberts et al., 2010; Pan et al., 2012; Lin et al., 2013). Similarly, convergent clinical studies have revealed a predictive longitudinal association between obesity and the development of age-related cognitive deficits (Cournot et al., 2006; Sabia et al., 2009; Dahl et al., 2013). Obesity is
also a risk factor for neurological diseases such as Alzheimer's disease (Frisardi et al., 2010; Farooqui et al., 2012). Interestingly, significant improvement in mood and cognitive function is reported after weight loss induced by bariatric surgery or diet restriction in obese subjects (Brinkworth et al., 2009; Andersen et al., 2010; Siervo et al., 2012; Alosco et al., 2014). Neuropsychiatric comorbidities considerably impair the quality of life and social functioning of obese individuals. More importantly, they emerge as additional risk factors for aggravation of obesity and related systemic pathological complications (Fiedorowicz et al., 2008; Scott et al., 2008). Indeed, lifetime history of neuropsychiatric disorders or stressful life events are reliable predictors for weight gain and the subsequent development of obesity (Goldbacher et al., 2009; Luppino et al., 2010; McIntyre et al., 2012). In addition, weight gain often appears as a common side effect of many psychiatric medications (Lopresti and Drummond, 2013). Altogether, these data point to a noxious bidirectional relationship between obesity and neuropsychiatric disorders. Identifying the pathophysiological mechanisms that underlie such a comorbid association represents therefore a major public health challenge.

Diet, social factors, and psychological distress are classically advanced as contributors to the development of neuropsychiatric symptoms in obese individuals. Mechanistically, these factors may impact the functioning of key biological systems participating in both obesity and neuropsychiatric disorders and able to target the central nervous system (CNS) (McIntyre et al., 2012; Hryhorczuk et al., 2013; Lopresti and Drummond, 2013). Although several biological components of obesity are likely candidates, there is increasing evidence for a role of inflammation in this process (Emery et al., 2007; Capuron et al., 2008, 2011a; Castanon et al., 2014; Lasselin and Capuron, 2014). Severe obesity is indeed associated with an inflammatory profile characterized by increased concentrations of circulating cytokines (Cannella and Clement, 2006). The effect of inflammation on brain function and behavior has been reported in many other chronic inflammatory conditions (Evans et al., 2005; Danzer et al., 2008). However, there remains a noticeable void in understanding the pathophysiological mechanisms underlying the development of neuropsychiatric symptoms in the context of obesity. We describe here how animal models of obesity, which already shed light on the mechanisms of weight control and associated metabolic alterations, can be useful to address this issue. We also review the available evidence showing that obesity is associated with exacerbated neuroinflammation leading to dysfunction in vulnerable brain regions associated with mood regulation, learning and memory, especially the hippocampus.

**Obesity: More than Just a Metabolic Disorder**

Obesity is defined by the World Health Organization as an “excessive or abnormal fat accumulation that presents a risk to health.” Weight gain is associated with marked hyperplasia of the white adipose tissue and substantial changes in the function of adipocytes that start to secrete a number of bioactive molecules collectively referred to as adipokines (leptin, adiponectin, and a multitude of pro- and anti-inflammatory cytokines) (Lehr et al., 2012; Aguilar-Valles et al., 2015). Due to their ability of acting remotely on different organs these molecules are suspected to participate in the etiology of many obesity-associated metabolic comorbidities (Lehr et al., 2012; Aguilar-Valles et al., 2015; Bluher and Mantzoros, 2015). Chronic obesity is indeed often associated with hypertension, coronary artery disease, dyslipidemia, hyperleptinemia, and impaired glucose tolerance linked to hyperinsulinemia and insulin resistance. Along with metabolic dysregulations, basal low-grade inflammation increasingly appears as another key component of obesity, which is now considered not only as a metabolic disorder but also as an inflammatory condition affecting both the innate and acquired immune systems (Schmidt and Duncan, 2003; Canella and Clement, 2006; Gregor and Hotamisligil, 2011). Elevated plasma levels of inflammatory cytokines (including interleukin (IL)-1β, tumor necrosis factor (TNF)-α, IL-6 and C-reactive protein) have been reported in obese patients (Park et al., 2005; Capuron et al., 2011a) and different animal models of obesity (Xu et al., 2003; De Souza et al., 2005; Cani et al., 2009; Pistell et al., 2010; Dinel et al., 2011, 2014; Lawrence et al., 2012). Interestingly, peripheral low-grade inflammation is significantly improved following weight loss induced by low-caloric diet or bariatric surgery in both obese humans (Ryan and Nicklas, 2004; Manco et al., 2007; Belza et al., 2009; Hakeem et al., 2009; Rao, 2012) and animals (Zhang et al., 2011; Liu et al., 2014; Schneck et al., 2014). Obesity is also characterized by increased susceptibility to immune-mediated diseases (Kanneganti and Dixit, 2012) and to infections (Amar et al., 2007; Rumel et al., 2010; Lawrence et al., 2012; Huttunen and Syrjanen, 2013; Dinel et al., 2014).

**Obesity: An Inflammatory Condition**

One major player in the development of the chronic low-grade inflammatory state characterizing obesity is the white adipose tissue, consistent with findings showing associations between circulating levels of cytokines and measures of central adiposity (Park et al., 2005). Together with adipocytes, infiltrated macrophages, and T cells that progressively accumulate in the adipose tissue potently secrete inflammatory mediators (Cannella and Clement, 2006; Gregor and Hotamisligil, 2011; Zeyda et al., 2011; Lasselin et al., 2013; Kim et al., 2014). In addition, accumulation of activated immune cells also releasing inflammatory factors in the circulation has been reported within other organs, in particular the liver and muscles (Pedersen and Febbraio, 2012; McNelis and Olefsky, 2014). More recently, gut microbiota alterations and increased gut permeability have also been involved in the pathogenesis of obesity and related metabolic comorbidities (Brun et al., 2007; Tehrani et al., 2012; Finelli et al., 2014), in particular through their impact on local and systemic inflammation (Cani et al., 2009, 2012; Verdam et al., 2013). Specific microbes-associated molecular patterns (MAMPs) such as gut microbiota-derived lipopolysaccharide (LPS) have been indeed shown to play a major role in the onset and progression of obesity-related inflammation and metabolic
diseases (Cani et al., 2007; Creely et al., 2007). Chronic intake of high-fat diet impairs gut permeability, which leads to the instauration of metabolic endotoxemia (i.e., increased plasma levels of LPS) contributing to obesity-related inflammation by activating systemic macrophages (Cani et al., 2008, 2009; Verdam et al., 2013). Conversely, reduced serum levels of an endotoxemia marker, the LPS binding protein, are found in obese individuals following weight loss (Cani et al., 2008; Yang et al., 2014).

Whatever the mechanisms triggering systemic inflammation in obesity, it is now clear that this inflammatory state contributes to increased central inflammatory processes (Rummel et al., 2010; Buckman et al., 2014) associated with both metabolic dysregulations and behavioral alterations (Dantzer et al., 2008). Obesity-related brain inflammation is particularly notable in the hypothalamus, with local enhanced inflammatory cytokine expression and activation of dependent signaling pathways being repeatedly reported in diet-induced obesity (DIO) models (De Souza et al., 2005; Zhang et al., 2008; Kleinridders et al., 2009; Cai and Liu, 2012; Gao et al., 2014; Maric et al., 2014). Increased infiltration, activation, and proliferation of microglia and astrocytes in the hypothalamus are also noted in both obese humans and rodent models of obesity (Thaler et al., 2012). Interestingly, increased hypothalamic inflammation has been related to the metabolic dysregulations characterizing severe obesity, including leptin resistance, insulin resistance and hyperglycemia (De Souza et al., 2005; Velloso et al., 2008; Zhang et al., 2008; Kleinridders et al., 2009).

Beyond basal hypothalamic inflammation that appears as a function of obesity, we also demonstrated in DIO models a marked exacerbation of hypothalamic expression of different inflammatory factors following a systemic immune challenge, namely acute intraperitoneal injection of LPS (Pohl et al., 2009; André et al., 2014). Of note, systemic LPS challenge also exacerbates expression of brain cytokines (IL-1β, TNF-α) in areas involved in mood regulation and memory formation such as the hippocampus (André et al., 2014; Boitard et al., 2014; Dinel et al., 2014). Besides, significant hippocampal inflammation is also reported in unstimulated conditions when more severe obesity is modeled (Dinel et al., 2011, 2014; Erion et al., 2014). For example, immune defects such as increased systemic inflammation and/or reduced immune competence (Cani et al., 2009; Rummel et al., 2010; Lawrence et al., 2012), which are reported in genetic models of severe obesity [ob/ob (deficient for leptin) and db/db (deficient for functional leptin receptor) mice], are associated with increased hippocampal cytokine (IL-1β, IL-6, TNF-α) expression (Dinel et al., 2011, 2014; Erion et al., 2014). Importantly, hippocampal inflammation is associated with signs of dysfunctions within the brain (Stranahan et al., 2008; Dey et al., 2014; Erion et al., 2014) and marked emotional and cognitive alterations (Stranahan et al., 2008; Dinel et al., 2011, 2014; Erion et al., 2014). These models of obesity are therefore especially suited for studying the long-term adverse effects of obesity and investigating in particular the molecular and cellular events underlying the development of emotional and cognitive alterations.

Cognitive and Emotional Alterations Displayed by Animal Models of Obesity

Several complementary models of obesity have been developed over the last decades in order to study obesity and related health complications, particularly cardiovascular diseases or type 2 diabetes (Varga et al., 2009; Kanasaki and Koya, 2011). Beyond these research topics, such animal models proved to be very useful for studying the neurobiological basis of obesity-associated emotional and cognitive alterations (Blessels and Gispen, 2005; Kanoski and Davidson, 2011). For example, models relying on diet modifications (DIO models), which are close to human obesity with respect to etiological aspects, give the opportunity of controlling the degree of obesity (low to moderate), as well as the duration and time of exposure to high-fat diet (perinatal periods, childhood, adulthood). Moreover, because of their longitudinal characteristic, DIO models allow for investigating the pathophysiological changes preceding the development of obesity-related comorbidities, including neuropsychiatric alterations. In addition, genetic models of obesity reproduce moderate to severe obesity and display most of the metabolic, inflammatory, and brain alterations characterizing this condition, including neurobehavioral alterations.

In agreement with clinical studies, experimentally-induced obesity is associated with a wide array of cognitive abnormalities including impairment in learning and memory (Kanoski and Davidson, 2011). For example, performances of spatial learning and long-term memory (Boitard et al., 2012, 2014; Valladolid-Acebes et al., 2013; André et al., 2014) or contextual fear conditioning (Hwang et al., 2010) are impaired in DIO models together with hippocampal synaptic plasticity (Molteni et al., 2002; Hwang et al., 2010; Boitard et al., 2012). Interestingly, the juvenile period (from weaning to young adulthood) seems to be particularly vulnerable to the adverse effects of high-fat diet on hippocampal function and related learning and memory processes (Valladolid-Acebes et al., 2013; Boitard et al., 2014). Indeed, exposure of adult mice to high-fat diet for the same duration as young mice does not yield such behavioral nor hippocampal alterations (Boitard et al., 2014; Valladolid-Acebes et al., 2013). Moreover, we recently showed that rats exposed to high-fat diet during the perinatal period and maintained on this diet during adulthood displayed impaired spatial memory and hippocampal neurogenesis in contrast with those exposed to the diet only during the perinatal period or after weaning (Lepinay et al., 2015). These data suggest therefore that exposure to high-fat diet during the perinatal period increases hippocampal vulnerability to the adverse effects of subsequent high-fat feeding. This study complements previously published data showing that maternal consumption of a high-fat diet can affect spatial memory (Bilbo and Tsang, 2010; Tozuka et al., 2010; Page et al., 2014) and hippocampal function in off-spring (Niculescu and Lupu, 2009; Tozuka et al., 2009). Similarly, an association between impaired hippocampus-dependent spatial memory performances in the water-maze or Y-maze tasks (Li et al., 2002; Dinel et al., 2011) and altered hippocampal neurogenesis, neuronal dendrite morphology, and
synaptic plasticity is also found in genetic models of obesity such as db/db mice (Stranahan et al., 2008, 2009; Erion et al., 2014; Ramos-Rodriguez et al., 2014). Interestingly, these mice also display increased hippocampal expression of cytokines that are known to influence hippocampal plasticity and behavior (Dinel et al., 2011). In addition, blockade of hippocampal IL-1β expression in db/db mice normalizes hippocampal dendritic spine density and prevents synaptic dysfunction and cognitive impairment (Erion et al., 2014). On the other hand, acquisition of a conditioned taste aversion learning task (Ohta et al., 2003), as well as working memory performances tested in a hippocampus-independent task are on the contrary preserved in db/db mice (Dinel et al., 2011). Altogether, these data point to the importance of the hippocampus as key brain area for mediating cognitive impairments linked to obesity.

Emotional reactivity is also impaired in animal models of obesity, although with a different time-course of development than cognition. Interestingly, we recently showed that DIO starting at weaning in mice alters first spatial memory, then anxiety-like behavior, whereas depressive-like behavior, either assessed in the forced swim test (FST) or tail suspension test (TST), remains unchanged unless the animals are challenged with LPS (André et al., 2014). Rats exposed to moderate high-fat diet from the perinatal period and throughout life also display memory impairments but unaltered depressive-like behavior in unstimulated conditions (Lepinay et al., 2015). Of note, increased basal depressive-like behavior has been previously reported in other DIO models but only when very high-fat diets associated with important metabolic dysfunctions and presumably basal low-grade inflammation are used (Yamada et al., 2011; Sharma and Fulton, 2013). Moreover, it is well-known that depressive-like behaviors mostly increase under challenging conditions such as stress exposure (Lu et al., 2008) or immune stimulation (Frenois et al., 2007; Moreau et al., 2008). Similarly, depressive-like behavior is also increased in obese mice compared to lean controls when experimental conditions used are particularly stressful (e.g., sustained and/or repeated exposure to the test, successive exposures to different behavioral tests over short periods of time) (Collin et al., 2000; Yamada et al., 2011). However, when stressful factors are tightly controlled, db/db mice display similar levels of depressive-like behavior than their lean db/+ controls, although anxiety-like behavior remains elevated (Dinel et al., 2011). These results strongly suggest an important role for the inflammatory system and/or the hypothalamo-pituitary-adrenal axis (HPA) axis, which are functionally related (Raison and Miller, 2003), in underlying emotional alterations associated with obesity. In addition, they indicate that obesity, in conjunction with environmental conditions, can amplify CNS dysfunctions and/or their harmful consequences on mood and cognition. Supporting this notion, it has been recently shown in DIO mice that exacerbated neuroinflammatory response to a systemic LPS challenge contributes to increase depressive-like behavior (Aguilar-Valles et al., 2014). Moreover, exacerbated neuroinflammation and depressive-like behavior displayed by LPS-injected DIO mice is also accompanied by enhanced HPA axis activation (André et al., 2014). Altogether, these data support the notion that obesity-associated cognitive and emotional alterations rely on interactions involving multiple systems, including metabolic characteristics, environmental influences and immune-related processes.

**Brain Inflammation: A Key Player in the Control of Cognitive Functions and Mood**

**From Sickness Behavior to Neuropsychiatric Symptoms**

Over the last decades, clinical (Evans et al., 2005; Raison et al., 2010; Capuron and Miller, 2011; Lasselin and Capuron, 2014) and experimental (Castanon et al., 2002; Frenois et al., 2007; Moreau et al., 2008) investigations focusing on the intricate relationship between the innate immune system and the brain have supported a main role for dysregulated production and/or brain action of cytokines in neuropsychiatric disorders through the profound action they exert on brain functions (Dantzer et al., 2008; Zunszain et al., 2012). Most cytokines have little or no function in healthy tissues, but they are rapidly induced locally by activated innate immune cells in response to tissue injury, infection, or inflammation. They are then able to act systemically on distant organs, including the brain, through a number of non-exclusive humoral, neural, and cellular pathways that allow the peripheral immune messages to be transmitted to the brain (Dantzer et al., 2008; Capuron and Miller, 2011). Activation of immune-to-brain communication ultimately induces the production of brain cytokines by activated endothelial and glial cells, particularly microglia (Layé et al., 1994; Castanon et al., 2004). Upon detection of homeostatic disturbances, microglia are transiently activated and rapidly engaged in brain adaptive immune responses mainly due to their ability to produce cytokines, express their receptors and amplify their signals (Ransohoff and Perry, 2009; Kettenmann et al., 2011). Within the brain, cytokines are able to influence pathways involved in behavioral regulations, including neurotransmitter metabolism and function, neuroendocrine activity, neural plasticity, and/or brain circuitry (Dantzer et al., 2008). During an infection, transient brain cytokine activation thus coordinate a large number of behavioral changes (including weakness, listlessness, malaise, anorexia, fatigue, and transient cognition and mood alterations) collectively referred to as sickness behavior. Necessary for infection recovery, sickness behavior usually resolves within few days, once microbial pathogens have been cleared and the innate immune system is no longer activated. However, failure to tightly regulate systemic immune activation and/or brain microglial activation leads to significant and prolonged induction of peripheral and brain cytokines. This induction in turn might culminate in medical conditions adversely affecting clinical outcomes, including neuropsychiatric symptoms, particularly when they ultimately affect key brain areas, such as the hippocampus, the cortex, or the amygdala (Dantzer et al., 2008). During the last decade, these findings have prompted a surge of interest for the circumstances precipitating the development of such pathological conditions and the identification of the underlying mechanisms.
Inflammation-associated Behavioral Alterations: Underlying Mechanisms

Converging clinical findings have shown that inflammation-related sickness behavior and neuropsychiatric symptoms differ by their respective duration and intensity, suggesting that they likely have distinct underlying mechanisms (Raison et al., 2010; Capuron and Miller, 2011). Cytokines are able to induce the synthesis of different enzymes in activated monocytes, macrophages and brain microglia, namely the indoleamine 2,3-dioxygenase (IDO) and GTP-cyclohydrolase 1 (GTP-CH1), which results in significant alterations in the biosynthesis of key monoamines (e.g., serotonin, dopamine) known to play a major role in mood regulation and cognitive function (Dantzer et al., 2008; Capuron et al., 2011b). As part of the immune response to infection, cytokine-induced activation of IDO, which is the first and rate-limiting enzyme degrading tryptophan along the kynurenine pathway, is usually beneficial to the host (Mellor and Munn, 2008). However, sustained brain IDO activation can also be deleterious by negatively impacting monoaminergic neurotransmission, but also neuronal survival. Interestingly, development of neuropsychiatric symptoms in medically ill patients chronically treated with IFN-α (Raison et al., 2010), elderly subjects (Capuron et al., 2011b), and patients suffering from Alzheimer's disease (Gulaj et al., 2010) or strokes (Gold et al., 2011) is associated with increased circulating levels of kynurenine. Severely obese individuals with high prevalence of neuropsychiatric comorbidity also display activation of IDO (Brandacher et al., 2006, 2007), together with larger hippocampal atrophy than non-obese subjects (Fotuhi et al., 2012). Increased brain kynurenine levels resulting from IDO activation can be further metabolized to produce neuroactive glutamatergic compounds, including 3-hydroxykynurenine (3HKyn) and quinolinic acid (QA), which play a key role in neuronal death and neurodegenerative diseases by stimulating NMDA receptors and promoting oxidative stress (Figure 1) (Stone et al., 2012; Campbell et al., 2014). On the other hand, kynurenine can also be metabolized in kynurenic acid (KA) that rather displays neuroprotective properties. However, these apparently antagonistic pathways are compartmentalized in the brain, with microglia preferentially producing QA, whereas astrocytes produce KA. Sustained immune activation therefore tips the scale in favor of neurotoxicity. In agreement with this finding, increased brain, or cerebrospinal fluid (CSF) concentrations of kynurenine and its neurotoxic metabolites have been reported in several neurodegenerative and psychiatric disorders (Schwarcz et al., 2001; Myint et al., 2007; Stone et al., 2012; Campbell et al., 2014). Moreover, they have been related with the stretch of brain damages, and with mood and cognitive impairments (Stone et al., 2012), suggesting that IDO activation may lead to both functional and structural alterations in the brain. Consistent with this assumption, activation of the kynurenine pathway has been recently shown to affect human hippocampal neurogenesis (Zunszain et al., 2012; Savitz et al., 2015a,b).

In line with clinical findings, experimental studies showed that development of depressive-like and anxiety-like behaviors induced by acute or chronic immune challenges in mice (Frenois et al., 2007; Godbout et al., 2008; Moreau et al., 2008; O’Connor et al., 2009a,b,c; Salazar et al., 2012; Corona et al., 2013; Lawson et al., 2013) is associated with increased peripheral and brain

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**FIGURE 1 | Tryptophan metabolism through the kynurenine pathway.**

Increased indoleamine 2,3-dioxygenase (IDO) activity occurring in activated monocytes, macrophages, and brain microglia in conditions of immune activation catabolizes tryptophan in kynurenine. Tryptophan is the biosynthetic precursor for the synthesis of serotonin. By reducing the availability of tryptophan for serotonin synthesis, IDO activation is able to impair serotonin neurotransmission. Kynurenine is metabolized in different neuroactive glutamatergic metabolites, including 3-hydroxykynurenine (3HKyn) and quinolinic acid that are produced by activated microglia, and kynurenic acid produced by astrocytes. Elevated levels of quinolinic acid are neurotoxic by activating glutamatergic NMDA receptors and promoting oxidative stress. High concentrations of kynurenic acid can be neuroprotective by antagonizing NMDA receptors, but sustained microglia activation rather promotes increased neurotoxicity. KAT, kynurenine aminotransferase; KMO, kynurenine monooxygenase; 3-HAO, 3-hydroxyanthranilic acid oxygenase; TPH, tryptophan hydroxylase.
IDO activity (Lestage et al., 2002; Moreau et al., 2005; André et al., 2008). Similarly, brain IDO activation was also associated with cognitive and emotional alterations following immune activation (Lawson et al., 2011; Barichello et al., 2013; Gibney et al., 2013; Xie et al., 2014). More importantly, pharmacological, or genetic inhibition of IDO activity prevents induction of depressive-like behaviors, anxiety-like behaviors and/or cognitive impairments by systemic immune challenges, without impacting sickness behavior (Henry et al., 2009; O’Connor et al., 2009a,b,c; Salazar et al., 2012; Barichello et al., 2013; Xie et al., 2014). In addition, aged mice (Godbout et al., 2008; Kelley et al., 2013), mice exhibiting constitutive microglial over-activation (Corona et al., 2010) and obese mice (André et al., 2014) display sustained cytokine production after an immune challenge, together with protracted brain IDO expression and depressive-like behavior (Godbout et al., 2008; Wynne et al., 2010; Corona et al., 2013). Further studies report that centrally induced inflammation, which only activates brain kynurenine pathway, is sufficient to elicit depressive-like behaviors (Fu et al., 2010; Park et al., 2011; Dobos et al., 2012; Lawson et al., 2013). In addition, other studies shed light on the hippocampus as important brain area for activation of cytokines and IDO (Frenois et al., 2007; André et al., 2008; Henry et al., 2009; Wang et al., 2009; Corona et al., 2010; Fu et al., 2010), although they are broadly stimulated within the brain in response to immune challenges (Castanon et al., 2004; André et al., 2008). Interestingly, emotional alterations linked to hippocampus IDO activation by an immune challenge are associated with reduced hippocampal expression of the brain-derived neurotrophic factor (BDNF) (Gibney et al., 2013). This neurotrophin contributes to mood regulation and memory function, including in conditions of immune activation (Barrientos et al., 2004), by supporting synaptic plasticity and neuronal excitability (Yamada and Nabeshima, 2003; Martinowich et al., 2007). Interestingly, cognitive impairment and emotional alterations reported in both obese DIO mice and db/db mice are also associated with reduced BDNF levels in the cortex (Pistell et al., 2010) and the hippocampus (Dinel et al., 2011). Reciprocally, interventions normalizing hippocampal levels of BDNF in these mice prevent hippocampus-mediated cognitive impairments (Moy and McNay, 2012; Kariharan et al., 2015). These results point to a link between increased neuroinflammation, impaired neurogenesis/synaptic plasticity, and behavioral alterations in obesity. Of note, brain IDO activation results in obese mice in a huge increase of brain kynurenine concentrations compared to lean mice, but similar impairment of brain tryptophan levels (André et al., 2014). These results support those showing that administration of kynurenine dose-dependently induces depressive-like behaviors, anxiety-like behaviors and cognitive impairment in normal-weight mice (Chess et al., 2009; O’Connor et al., 2009c; Alexander et al., 2012; Salazar et al., 2012; Agudelo et al., 2014). Altogether, although these results do not exclude the possible role of impaired serotonin synthesis in inflammation-induced neuropsychiatric symptoms, they clearly support a key role for the neuroactive kynurenine metabolites resulting from IDO activation.

Concurrently with activation of IDO, chronic inflammation also activates GTP-CH1, which is the rate limiting enzyme of GTP conversion ultimately leading to the production of neopterin by activated immune cells at the expense of tetrahydrobiopterin (BH4) formation (Figure 2) (Murr et al., 2002; Oxenkrug, 2010; Capuron et al., 2011b). By acting as a cofactor of aromatic amino acid hydroxylase, BH4 plays a fundamental role in neurotransmitter synthesis, including serotonin (Capuron et al., 2011b) but mainly dopamine (Neurauter et al., 2008). Cytokine-induced GTP-CH1 activation, classically assessed by measuring increased production of neopterin, is indeed able to impair the dopaminergic neurotransmission (Felger and Miller, 2012) which is known to be involved in mood disorders and cognitive dysfunctions, including in conditions of chronic immune stimulation (Brydon et al., 2008; Capuron et al., 2012). Consistent with these findings, reduced BH4 levels have been reported in patients with psychiatric disorders (Hashimoto et al., 1994; Hoekstra et al., 2001). Similarly, increased blood neopterin concentrations correlate with a greater incidence of depressive episodes in patients with major depression (Celik et al., 2010). Interestingly, chronic low-grade inflammation that is classically reported in elderly people is associated with activation of both IDO and GTP-CH1 (Oxenkrug, 2010; Capuron et al., 2011b). More importantly, these enzymatic activations have been shown to participate in the pathophysiology of neuropsychiatric symptoms, whose prevalence is often high in aged population (Oxenkrug, 2010; Capuron et al., 2011b). Of note, in addition to impair monoaminergic neurotransmission, both enzymes contributes to increase oxidative stress (Felger and Miller, 2012; Stone et al., 2012; Campbell et al., 2014). Increased neopterin levels have been reported in obese rats (Finkelstein et al., 1982) and humans (Ledochowski et al., 1999; Oxenkrug et al., 2011), suggesting that activation of the GTP-CH1 enzyme by cytokines, and the consequent alterations of dopamine neurotransmission, may contribute to the development of neuropsychiatric symptoms reported in obesity. Although this assumption still needs to be confirmed, it is worth mentioning that alterations of dopamine function, together with alterations in basal ganglia/reward circuitry have been reported in obese patients (de Weijer et al., 2011; Volkow et al., 2011). Moreover, depressive-like behavior is associated in DIO mice with alterations in striatal circuitry, supporting a role for dopamine-related disruptions in obesity-associated depressive symptoms (Sharma and Fulton, 2013).

Factors Modulating the Impact of Neuroinflammation on Cognitive and Emotional Alterations Associated with Obesity

HPA Axis

Brain effects of cytokines on mood regulation and cognitive function are likely modulated by the tight interactions existing between the inflammatory and neuroendocrine systems, in particular the HPA axis (Raison and Miller,
FIGURE 2 | Effect of immune activation on the availability of tetrahydrobiopterin (BH4). The guanosine-triphosphate-cyclohydrolase-1 (GTP-CH1) is the rate limiting enzyme of guanosine-triphosphate (GTP) conversion in dihydrobiopterin (BH2). Increased GTP-CH1 activity occurring in conditions of immune activation ultimately leads to the production of neopterin at the expense of tetrahydrobiopterin (BH4) formation. BH4 plays a fundamental role in neurotransmitter synthesis, in particular serotonin and dopamine, by acting as a cofactor of the tryptophan hydroxylase (TPH) and tyrosine hydroxylase (TH). Cytokine-induced GTP-CH1 activation reduces therefore monoaminergic neurotransmission. BH4 is also a cofactor of the nitric oxide synthase (NOS). Reduced BH4 availability can indirectly contribute to increase the production of free radicals promoting oxidative stress.

2003; Capuron and Miller, 2011) which is highly activated in obesity (Dinel et al., 2011, 2014). Immune alterations are indeed notorious for causing significant changes in HPA axis activity and vice versa. Consequently, alterations of HPA axis functions and inflammatory activation are often associated in many pathological conditions, including cancers treated by immunotherapy (Capuron et al., 2003), mood disorders (Zunszain et al., 2011) or obesity (Dinel et al., 2011, 2014; André et al., 2014). Interestingly, glucocorticoids have been recently shown to sensitize microglia in an animal model of obesity (Dey et al., 2014). Reciprocally, DIO mice display exacerbated HPA axis activation in response to an immune challenge, together with increased neuroinflammation and depressive-like behavior (André et al., 2014). This result supports the notion that inflammatory factors and glucocorticoids may act together in the context of obesity to promote the development of depressive symptoms (Hryhorczuk et al., 2013) and cognitive functions, in particular those involving the hippocampus (Stranahan et al., 2008; Dey et al., 2014). Interestingly, both cytokines and glucocorticoids have been shown for example to impair hippocampal neurogenesis and neuronal function in animal models of obesity (Stranahan et al., 2008; Dinel et al., 2011; Erion et al., 2014; Wosiski-Kuhn et al., 2014). Moreover, reversing the impairment of hippocampal neurogenesis by targeting cytokines or glucocorticoids improves spatial memory deficits displayed by obese mice (Stranahan et al., 2008; Erion et al., 2014).

Leptin
One of the stronger features associated with obesity is the significant increase of leptin production (Lehr et al., 2012). This adipokine has been extensively studied for decades for its key role in the control of energy homeostasis and feeding behavior through activation of specific leptin receptors in the hypothalamus (Rosenbaum and Leibel, 2014). However, due to the broad brain distribution of leptin receptors, which are also located throughout the cortex and the hippocampus, leptin has been shown to modulate memory processes and mood disorders (Guo et al., 2013; Farr et al., 2015). Leptin is also known to locally facilitate synaptic plasticity and neurogenesis (Shanley et al., 2001). Deficits in spatial memory reported in DIO mice occur concomitantly with desensitization of leptin signaling pathway in the hippocampus (Valladolid-Acebes et al., 2013). Moreover, selective deletion of leptin receptors in adult hippocampus induces depressive-like behavior in mice (Guo et al., 2013) and confers resistance to behavioral antidepressant effect of fluoxetine (Guo and Lu, 2014). In addition, obese mice which are characterized by impaired leptin signaling pathway (e.g., db/db or ob/ob mice) display deficits of hippocampal-dependent memory and increased anxiety-like behavior (Stranahan et al., 2008; Dinel et al., 2011), together with increased basal hippocampal inflammation (Dinel et al., 2011; Erion et al., 2014) and altered neuroinflammatory response to an immune challenge (Rummel et al., 2010; Lawrence et al., 2012; Dinel et al., 2014). These
findings suggest that both leptin and cytokines may contribute together to the development of behavioral alterations associated with obesity. Interestingly, this assumption is supported by mounting evidence showing that leptin acts as an important neuroendocrine modulator of the immune system by modulating the activation of both peripheral immune cells and brain microglia (Lafrance et al., 2010; Aguilar-Valles et al., 2015).

**Insulin**

This hormone, whose circulating levels and signaling pathway are altered in obesity, is able to interact with inflammatory processes and to act within the brain to modulate mood and cognition (Ghasemi et al., 2013). Impaired insulin signaling pathway may therefore, as with leptin, contribute to the development of neuropsychiatric symptoms in the context of obesity by interacting with brain cytokines. More studies are necessary to test this hypothesis. However, it is worth noting that several studies including ours suggest that insulin *per se* is not essential or sufficient to explain behavioral alterations occurring in obesity. For example, the increased risk of cognitive dysfunction displayed by patients with metabolic syndrome is independent from the presence of diabetes (Muller et al., 2010). Consistent with these clinical findings, we recently reported increased emotional behaviors and cognitive impairment in DIO animals in the absence of any significant hyperinsulinemia (André et al., 2014; Boitard et al., 2014). Similarly, normalization of peripheral hyperglycemia in *db/db* mice does not improve spatial cognitive impairments or anxiety-like behaviors (Stranahan et al., 2008, 2009). Besides, this normalization does not alter brain concentrations of glucose and insulin that are similar in both *db/db* and *db/+* mice (Stranahan et al., 2008).

**Gut Microbiota**

Converging evidence shows that obesity-related microbiota dysregulations, which play a critical role in induction of peripheral and brain inflammation (Bruce-Keller et al., 2015), impact mood and cognitive functions (Cryan and Dinan, 2012), although the nature of the biological pathways (neuronal, hormonal and/or immune) underlying this effect is still elusive (Cryan and Dinan, 2012). Interestingly, it has been recently

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**FIGURE 3 | Proposed role of neuroinflammation in obesity-associated cognitive and emotional alterations.** Obesity is characterized by metabolic alterations (hyperinsulinemia, hyperleptinemia, increased activation of the hypothalamo-pituitary-adrenal (HPA) axis... and peripheral low-grade inflammation, which originates from alterations in adipose tissue and gut functions. These obesity-associated alterations are well-known to promote brain inflammatory processes that represent key players in the development of behavioral alterations associated with obesity. By sustaining neuroinflammation, as manifested by chronic activation of microglia, brain production of inflammatory cytokines, and local activation of indoleamine 2,3-dioxygenase (IDO) and guanosine-triphosphate-cyclohydrolase-1 (GTP-CH1), obesity may impair monoaminergic neurotransmission, neurogenesis, and synaptic plasticity, and concomitantly promote neurotoxicity. Such alterations of brain function induced by neuroinflammation likely represent major pathophysiological pathways to cognitive and emotional alterations in obesity.
reported that transplantation of gut microbiota from DIO mice to lean mice is sufficient to induce both brain microglial activation and neurobehavioral changes in the absence of obesity (Bruce-Keller et al., 2015). This elegant study supports the notion that obesity-related gut microbiota alterations may modulate gut-to-brain communication pathways, leading to the development of neuropsychiatric comorbidities associated with neuroinflammation. Akin with this assumption, the use of compounds which beneficially alter the microbiota (e.g., prebiotics or probiotics) appears as a promising way to improve neuropsychiatric comorbidities in obese patients (Cryan and Dinan, 2012). More generally, nutritional interventions based on factors with immunomodulatory properties and known impact on behavior and mood, in particular omega-3 polyunsaturated fatty acids and antioxidants (see Gomez-Pinilla and Nguyen, 2012; Bazinet and Layé, 2014 for review), are tractable strategies for developing novel therapeutics for obesity-related neuropsychiatric disorders. Lastly, based on the key role of the kynurenine pathway in altering mood and cognition (Dantzer et al., 2008), the opportunity of directly targeting this pathway, as promisingly tested in the context of cancer (Platten et al., 2015), should likely represent another interesting therapeutic approach.

Conclusion

Altogether, data presented in this review clearly show that brain inflammatory processes represent key players in the development of neuropsychiatric comorbidities in obesity. These likely rely on interactions involving multiple systems, including inflammatory processes but also neuroendocrine systems, gut microbiota and environmental influences. Indeed, neuroinflammation appears to be the cornerstone of the different factors contributing to induce neuropsychiatric symptoms in obesity. Several issues are however still unclear, in particular the identification of the specific brain pathways and/or mechanisms targeted by neuroinflammation and underlying obesity-related mood and cognitive alterations. Recent experimental results reported in this review suggest that cytokine-induced unbalanced relationship between neurogenesis and neuronal death combined with alterations in monoamine metabolism and function likely represent a major pathophysiological pathway to neuropsychiatric comorbidities in obesity (Figure 3). Given the alarming and continuous rise in obesity in modern societies and its role as a risk factor for many other diseases including neuropsychiatric disorders, therapeutic strategies to reduce obesity-related inflammation may be very promising approach to improve the quality of life and health outcomes of overweight and obese individuals.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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