Review: brain neurobiology of gambling disorder based on rodent models

Abstract: Different literature reviews of gambling disorder (GD) neurobiology have been focused on human studies, others have focused on rodents, and others combined human and rodent studies. The main question of this review was: which are the main neurotransmitters systems and brain structures relevant for GD based on recent rodent studies? This work aims to review the experimental findings regarding the rodent’s neurobiology of GD. A search in the Pub Med database was set (October 2012–October 2017) and 162 references were obtained. After screening, 121 references were excluded, and only 41 references remained from the initial output. More, other 25 references were added to complement (introduction section, neuroanatomical descriptions) the principal part of the work. At the end, a total of 66 references remained for the review. The main conclusions are: 1) according to studies that used noninvasive methods for drug administration, some of the neurotransmitters and receptors involved in behaviors related to GD are: muscarinic, N-methyl-D-aspartate (NMDA), cannabinoid receptor 1 (CB1), cannabinoid receptor 2 (CB2), dopamine 2 receptor (D2), dopamine 3 receptor (D3), and dopamine 4 receptor (D4); 2) moreover, there are other neurotransmitters and receptors involved in GD based on studies that use invasive methods of drug administration (eg, brain microinjection); example of these are: serotonin 1A receptor (5-HT1A), noradrenaline receptors, gamma-aminobutyric acid receptor A (GABAA), and gamma-aminobutyric acid receptor B (GABAB); 3) different brain structures are relevant to behaviors linked to GD, like: amygdala (including basolateral amygdala (BLA)), anterior cingulate cortex (ACC), hippocampus, infralimbic area, insular cortex (anterior and rostral agranular), nucleus accumbens (NAc), olfactory tubercle (island of Calleja), orbitofrontal cortex (OFC), medial prefrontal cortex (mPFC), prefrontal cortex (PFC) – subcortical network, striatum (ventral) and the subthalamic nucleus (STN); and 4) the search for GD treatments should consider this diversity of receptor/neurotransmitter systems and brain areas.

Keywords: gambling disorder, review, nervous system, murine paradigm

Introduction

Literature reviews regarding gambling disorder (GD) neurobiology have been specialized on human, rodents, or combination of both. The main question of this review was: which are the main neurotransmitters systems and brain structures relevant for GD based on recent rodent studies? Let me define first gambling and its epidemiological traits, before going in its neurobiological aspects.

Gambling conduct can be described as to put in peril anything significant, and to confide on the assumption of obtaining a gain in return. GD is characterized by gaming behaviors that seriously disrupt the finances, social relations, and professional advancement of a fellow. The lifetime prevalence of GD has been estimated
Moreover, GD is presently included in the Diagnostic and Statistical Manual of Mental Disorders-5 (DSM)-5, in a novel category, within the division of addictions (behavioral addictions). There have been some literature reviews of the neuropsychobiology of GD focused on human clinical studies. Specifically, the Lemieux and al’Absi’s review proposed that psychological and neurobiological aspects of the stress play a significant role in the starting, prolongation, and relapse of the addictions (including GD). Moreover, the mechanisms include interactions between biological mediators of the stress and the reward system; also, interactions between mediators of the stress and other systems related to addiction (endogenous opioids, the sympathetic-adrenal-medullary system, and endocannabinoids).

Another review work by Grant et al posed that GD is linked with alteration across different cognitive domains related to impulsivity and compulsivity; moreover, it pointed that, based on imaging reports, GD relates to anatomical and functional anomalies of nexus involved in the reward processing and top-down monitoring. In addition, it pointed that probably, diverse neural systems are involved in the pathophysiology (related to serotonin (5-HT), glutamate, dopamine (DA), opioids, and norepinephrine). Then again, the Goulet-Kennedy et al’s review points that prefrontal cortex (PFC) and the striatum are the main conductors of decision processes, based on clinical studies. Furthermore, that literature review states that the traits of decision making’s neural networks can be characterized by means of imaging technology; also, they consider that non-invasive neural stimulation in the PFC, and its network (striatum and others) have elucidated the neurobiological basis of decision-making processes; Decision making is involved in different aspects of our daily life, including all the spectrums of gambling behavior. Hence, a better understanding of the decision process’ neurobiology could be useful for a better quality of life of patients.

The publications review of the Banz et al group emphasizes in the capacity of neurobiological data to help in the promotion of improved norms and strategies for treatment and prevention. Furthermore, another review by Levy and Glimcher concludes that imaging investigations in humans suggest the existence of a brain network that codifies the values of rewards by means of a standard neural scale. Based on the authors, the brain area linked with this standard neural scale is a zone of the ventromedial prefrontal cortex (vmPFC)/orbitofrontal cortex (OFC). The authors propose that a better comprehension of brain mechanism for estimating and deciding might provide basic discernments of abnormal choice conducts like those of gambling.

Also, an imaging meta-analyses review by Meng et al reports that GD fellows display a significantly higher activation (compared to healthy controls) in brain areas like right lentiform nucleus and left middle occipital gyrus. Moreover, the South Oaks Gambling Screen scores were linked with overactivity in the right lentiform nucleus and in the bilateral parahippocampus; but the scores were negatively linked to right middle frontal gyrus. Altogether, this suggests dysfunction within the frontostriatal cortical pathway in fellows with GD.

In addition, other reviews focus on both human and rodent. The Norbury and Husain publications review points that a high level of sensation seeking is a factor related to gambling and substance addiction. Moreover, these authors support the existence of a relationship between sensation seeking and dopaminergic transmission, especially in the D2 receptors. Specifically, fellows with marked sensation seeking display also elevated DA tonic levels and an over-responsive midbrain dopaminergic responses to signals of future reward. Moreover, Norbury and Husain propose that even for stimuli of similar strength, reactive responses could vary in terms of approach-avoidance displayed by the subject; the authors propose that these variations stem from differences in the efficiency of DA transmission at the level of the striatum.

Additionally, another review by Quintero concludes that pathological and nonpathological gamblers can differ in terms of brain’s anatomy, brain’s physiology, electroencephalography (EEG) profile, executive and cognitive efficiency. For instance, fellows with GD can denote alterations in the insula, OFC, and frontal lobe; moreover, fellows with GD compared to nonpathological gambler show differences in frontoparietal activation pattern (if winning or losing a game) and insular activity (altered cognitive interpretation of near-miss results and trial success) related to gaming. With respect to anatomical differences between gamblers and non-gamblers, the first ones show more gray-matter volume compared to normal subjects, based on magnetic resonance imaging technology; furthermore, gamblers have a smaller size of right thalamus, right hippocampus, and left putamen compared to normal subjects. Regarding research on
Another review by Anselme stresses that the correctness of gambling decision is affected by the action of DA receptors and brain areas like insular cortex (rostral agranular zone), infralimbic, and prelimbic.6

Another review by Potenza stresses that diverse neurotransmitters like glutamate, noradrenaline, DA, 5-HT, opioid, and brain structures like insula, ventral striatum, and vmPFC (among other areas) are linked to gambling and GD.15 Furthermore, a literature review by van den Bos et al did focus on three almost ignored aspects of GD: developmental sex differences in GD, adolescence as a sensitive period for developing GD, and paths for upgrading ecological validity of investigative tools.16

Finally, another set of reviews has been specialized on rodents.3,4,17,18 Particularly, a literature review by Alguacil and González-Martin stressed on the “umbrella category” of reward deficiency syndrome; this syndrome includes diverse neuropsychiatric and addiction disorders (including gambling).3 More, these disorders share dysfunctional reward sensitivity, inadequate impulsivity, and/or compulsive conduct. That review considers that further investigation about the reward deficiency syndrome could ease the design of new drugs that are efficient for that cluster of disorders.3

A publications review by Winstanley and Clark emphasizes that the adequate laboratory models for GD should screen fundamental cognitive procedures and have adequate translatability to different species;4 moreover, models with these characteristics have a potential capacity to contribute to decision/neuroscience and the investigation of addictive behaviors.4 Another review by Anselme stresses that deprivation and randomness, either psychological or physiological, increase the motivation for searching valuable stimuli;17 moreover, this increase in motivation relates to the organism’s hardness for forecasting relevant environment’s stimulus and incidents.17

Finally, a review by Cocker and Winstanley states that cognitive biases are important in the evolution of GD;18 furthermore, these biases can be recreated in rodent models. In effect, some evidences suggest that biases can be linked to dopaminergic activity, especially in the D₄ receptor; the authors suggest the exploration of the D₄ receptor as an alternative for treating GD.18

The present review aims to integrate recent research findings about the rodent neurobiology of GD and related behaviors. It is expected that this integration could ease the elaboration of most complete pharmacological and/or behavioral approaches for treating GD in human populations.

Materials and methods
Inclusion and exclusion criteria
The publications were selected based on the next inclusion criteria: a) rodent studies (mice or rat), b) experimental or quasi-experimental design, c) publications that include description about the relationship between the nervous system (brain and/or neurotransmitter) and gambling behavior or GD, d) publications that detail the number of animals, e) the sex of the animals could be male, female or not specified in the publication, f) publications written in English (at least title and abstract), and g) publications released within a recent five years temporal range: October 2012–October 2017. As a reference, some reviews were added, but principally for the introduction and discussion parts.

With respect to the exclusion criteria of the publications, these included the next: a) the non-compliance of the inclusion criteria, and b) it should not be an abstract, nor a publication of a scientific meeting, nor a publication included in non-scientific literature.

Inquiry strategy
A screening of publications in the Pub Med database was carried out based on the five recent years (October/01/2012–October/20/2017). The search terms included: “Gambling” AND “Brain”, “Gambling” AND “Neurobiology”, “Gambling Disorder” AND “Brain”, and “Gambling Disorder” AND “Neurobiology”. The next filters were added for the searching process: text availability (Abstract), species (other animals), Languages (English). Initially, 162 references were obtained in the Pub med search. A total of 121 references was eliminated by different factors (literature review or meta-analysis type, non-English language, human specie, duplicates, and others), resulting in a total of 41 references for subsequent analysis. In addition, another 25 complementary references were detected through references scanning or web searching, and added to the manuscript. As a reference, these 25 references related mainly to the background in the field (included in the introduction section) and to neuroanatomical references. The complete (full) forms of these 66 references were obtained on the web or solicited directly to the authors; later these references were evaluated for the preparation of the review.

More information is described in Figure 1 (Flow diagram of publication selection process). As a guide for the
Results

There are two tables summarizing the results: Table 1 entitled “Summary of main studies included in the review – Neurotransmitters”, and Table 2 entitled “Summary of main studies included in the review – Brain structures”. These tables detail different aspects of the studies revised like: neurotransmitters, messengers and receptors systems studied, the brain area, drug name, drug effects, drug route of administration, mental process and/or conduct analyzed, behavioral test (paradigm) used,
Table 1 Summary of main studies included in the review – Neurotransmitters

<table>
<thead>
<tr>
<th>Author &amp; year</th>
<th>Messenger &amp; receptor</th>
<th>Brain area</th>
<th>Drug name, effect, route</th>
<th>Mental process and/or conduct analyzed</th>
<th>Behavioral test</th>
<th>Species</th>
<th>Relevance of study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silveira et al, 2015</td>
<td>Acetylcholine – muscarinic</td>
<td>na</td>
<td>Scopolamine, antagonist, sc</td>
<td>Decision making</td>
<td>Rodent gambling task</td>
<td>Rats</td>
<td>Receptor affect decision making under conditions of risk and uncertainty</td>
</tr>
<tr>
<td>Yates et al, 2015</td>
<td>Glutamate – NMDA</td>
<td>na</td>
<td>MK-801, antagonist, sc</td>
<td>Decreased sensitivity to reinforced amount without altering delay/probability discounting</td>
<td>Delayed/probabilistic reinforcement, sensitivity to reinforcer amount</td>
<td>Rats</td>
<td>- NMDA differ mediating impulsivity</td>
</tr>
<tr>
<td>Yates et al, 2015</td>
<td>Glutamate – NMDA</td>
<td>na</td>
<td>Ketamine, antagonist, ip</td>
<td>Decreased sensitivity to reinforcer amount</td>
<td>Delayed/probabilistic reinforcement, sensitivity to reinforcer amount</td>
<td>Rats</td>
<td>- NMDA differ mediating impulsivity</td>
</tr>
<tr>
<td>Gueye et al, 2016</td>
<td>Cannabinoids – CB1/CB2 receptors</td>
<td>na</td>
<td>WIN 55, 212–2, agonist, ip</td>
<td>Improved choice strategy and increased choice latency (SO group); but increased perseverations punished (O group)</td>
<td>Rodent gambling task</td>
<td>Rats</td>
<td>Agonism of CB1/CB2 affect gambling choice differentially in healthy and dysfunctional subjects</td>
</tr>
<tr>
<td>Di Ciano et al, 2015</td>
<td>Dopamine – D4 receptor</td>
<td>na</td>
<td>PD 168077, agonist, ip</td>
<td>- D4 agonist small decrease latencies</td>
<td>Rodent gambling task</td>
<td>Rats</td>
<td>Modest effect on latency in decision making</td>
</tr>
<tr>
<td>Pes et al, 2017</td>
<td>Dopamine – D2/D3 receptor</td>
<td>na</td>
<td>Pramipexole, agonist, sc</td>
<td>Increase unfavorable decisions, impaired the discounting of probabilistic losses, and augment risk-taking behaviors</td>
<td>Probability discounting task</td>
<td>Rats</td>
<td>- Pramipexole distorted representational rewards and impaired ability to discern contingencies (unfavorable and favorable) - Pramipexole conduct effects dissociates from changes in mesolimbic dopamine</td>
</tr>
</tbody>
</table>

Abbreviations: CB1, cannabinoid receptor 1; CB2, cannabinoid receptor 2; D3, dopamine 2 receptor; D4, dopamine 4 receptor; GD, gambling disorder; ip, intraperitoneal administration; NMDA, N-methyl-D-aspartate; na, not applicable (because non-specific brain area was studied); O group, optimal group; sc, subcutaneous administration; SO group, suboptimal group.
Table 2 Summary of main studies included in the review – Brain structures

<table>
<thead>
<tr>
<th>Author &amp; year</th>
<th>Brain area</th>
<th>Messenger &amp; receptor</th>
<th>Drug name, effect, route</th>
<th>Behavioral test</th>
<th>Other technique/treatment used</th>
<th>Mental process and/or conduct analyzed</th>
<th>Species</th>
<th>Relevance of study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fitoussi et al, 2015</td>
<td>Amygdala</td>
<td>Serotonin, Serotonergic receptor</td>
<td>—</td>
<td>Rat gambling task</td>
<td>Lesion of basolateral amygdala</td>
<td>Decision making</td>
<td>Rat</td>
<td>Amygdala: low serotonergic metabolism and constant action related to poor decision</td>
</tr>
<tr>
<td>Tremblay et al, 2014</td>
<td>Basolateral amygdala</td>
<td>—</td>
<td>—</td>
<td>Loss – chasing task, the betting task</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Zeeb &amp; Winstanley, 2013</td>
<td>Orbitofrontal cortex – basolateral amygdala pathway</td>
<td>—</td>
<td>—</td>
<td>Rat gambling task</td>
<td>Disconnection of pathway</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Cocker et al, 2016</td>
<td>Anterior cingulate cortex</td>
<td>GABA, GABA&lt;sub&gt;A&lt;/sub&gt; and GABA&lt;sub&gt;B&lt;/sub&gt;</td>
<td>Muscimol, agonist, infusion and baclofen, agonist, infusion</td>
<td>PD168077, agonist, infusion</td>
<td>Rodent slot machine task</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Cocker et al, 2016</td>
<td>Cingulate cortex (anterior)</td>
<td>—</td>
<td>—</td>
<td>Rat</td>
<td>Rat</td>
<td>Qualitatively equal effect (compared to left column – GABA), but augmentation reward expectancy was only evident on archetypal &quot;near-miss&quot; trials</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Pittaras et al, 2016</td>
<td>Hippocampus</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Quintero Garzola</td>
<td></td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

(Continued)
<table>
<thead>
<tr>
<th>Author &amp; year</th>
<th>Brain area</th>
<th>Messenger &amp; receptor</th>
<th>Drug name, effect, route</th>
<th>Behavioral test</th>
<th>Other technique/treatment used</th>
<th>Mental process and/or conduct analyzed</th>
<th>Species</th>
<th>Relevance of study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Koot et al, 2013</td>
<td>Insular cortex</td>
<td>Corticosteroid hormones</td>
<td>—</td>
<td>Rodent gambling task</td>
<td>c-fos immunohistochemistry</td>
<td>Decision making</td>
<td>Rat</td>
<td>Stress (corticosteroid rapid action) disrupts decision making (reward-based)</td>
</tr>
<tr>
<td>Mizoguchi et al, 2015</td>
<td>Insular cortex</td>
<td>GABA, GABA&lt;sub&gt;AL&lt;/sub&gt; and GABA&lt;sub&gt;B&lt;/sub&gt;</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>Rat</td>
<td>Insular cortex inactivation induces risk-taking behaviors linked to altered decision-making during tests</td>
</tr>
<tr>
<td>Ishii et al, 2012</td>
<td>Anterior insular cortex</td>
<td>GABA, GABA&lt;sub&gt;AL&lt;/sub&gt; and GABA&lt;sub&gt;B&lt;/sub&gt;</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>Risk taking behaviors</td>
<td>Rat</td>
<td>Anterior insular cortex inactivation decreased risk preference</td>
</tr>
<tr>
<td>Ishii et al, 2015</td>
<td>Anterior insular cortex</td>
<td>Dopamine, D&lt;sub&gt;2&lt;/sub&gt;</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>Risk preference</td>
<td>Rat</td>
<td>Anterior insular cortex blockade increased risk preference, after winning in a previous risky choice</td>
</tr>
<tr>
<td>Ishii et al, 2015</td>
<td>Anterior insular cortex</td>
<td>Serotonin, 5-HT&lt;sub&gt;1A&lt;/sub&gt;</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>Risk preference</td>
<td>Rat</td>
<td>Anterior insular cortex blockade increased risk preference, after losing in a previous risky choice</td>
</tr>
<tr>
<td>Pushparaj et al, 2015</td>
<td>Caudal agranular Insular cortex</td>
<td>Muscimol, agonist, infusion and Baclofen, agonist, infusion</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>Rat</td>
<td>Caudal agranular Insular cortex is not involved in decision-making behavior under risk</td>
</tr>
</tbody>
</table>

(Continued)
<table>
<thead>
<tr>
<th>Author &amp; year</th>
<th>Brain area</th>
<th>Messenger &amp; receptor</th>
<th>Drug name, effect, route</th>
<th>Behavioral test</th>
<th>Other technique/treatment used</th>
<th>Mental process and/or conduct analyzed</th>
<th>Species</th>
<th>Relevance of study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pushparaj et al, 2015</td>
<td>Rostral agranular insular cortex</td>
<td>GABA, GABA_A, and GABA_B</td>
<td>Muscimol, agonist, infusion and baclofen, agonist, infusion</td>
<td>Rodent gambling task</td>
<td>Lesion by means of ibotenic acid infusion</td>
<td>Decision making: favor options with high reward frequency and low punishment</td>
<td>Rat</td>
<td>Rostral agranular insular cortex is involved in decision-making behavior under risk</td>
</tr>
<tr>
<td>Peak et al, 2015</td>
<td>Lateral ventricle</td>
<td></td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>Decision making</td>
<td>Rat</td>
<td>Lateral ventricles enlargement did not alter decision making of the offspring</td>
</tr>
<tr>
<td>Fitoussi et al, 2015</td>
<td>Infrafimbic area</td>
<td>Serotonin</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>Decision making</td>
<td>Rat</td>
<td>Infrafimbic higher serotonergic metabolism relates to poor decision making</td>
</tr>
<tr>
<td>Koot et al, 2013</td>
<td>Infrafimbic area</td>
<td>Corticosteroid hormones</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>Decision making</td>
<td>Rat</td>
<td>Stress (corticosteroid rapid action) disrupts decision making (reward-based)</td>
</tr>
<tr>
<td>Zeeb et al, 2015</td>
<td>Infrafimbic area</td>
<td>GABA, GABA_A, and GABA_B</td>
<td>Muscimol, agonist, infusion and baclofen, agonist, infusion</td>
<td>Rodent gambling task</td>
<td>---</td>
<td>Decision making (disrupted by inactivation)</td>
<td>Rat</td>
<td>Inactivation of infrafimbic area has effects on option choice: increase for bad and decrease for good choices</td>
</tr>
<tr>
<td>Zeeb et al, 2015</td>
<td>Infrafimbic area</td>
<td>Dopamine, D_2</td>
<td>Eticlopride hydrochloride, antagonist, infusion</td>
<td>Rodent gambling task</td>
<td>---</td>
<td>Choice preference (not altered)</td>
<td>Rat</td>
<td>Infrafimbic disruption does not alter choice preference nor optimal performance</td>
</tr>
</tbody>
</table>

(Continued)
Table 2 (Continued)

<table>
<thead>
<tr>
<th>Author &amp; year</th>
<th>Brain area</th>
<th>Messenger &amp; receptor</th>
<th>Drug name, effect, route</th>
<th>Behavioral test</th>
<th>Other technique/treatment used</th>
<th>Mental process and/or conduct analyzed</th>
<th>Species</th>
<th>Relevance of study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zeeb et al, 2015</td>
<td>Prelimbic cortex</td>
<td>GABA, GABA&lt;sub&gt;A&lt;/sub&gt;, and GABA&lt;sub&gt;B&lt;/sub&gt;</td>
<td>Muscimol, agonist, infusion and baclofen, agonist, infusion</td>
<td>Rodent gambling task</td>
<td>Maternal vitamin D depletion (before and during pregnancy)</td>
<td>Choice selection was unaffected by inactivation</td>
<td>Rat</td>
<td>Prelimbic inactivation: does not alter choice preference</td>
</tr>
<tr>
<td>Peak et al, 2015</td>
<td>Neocortex</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>Electrophysiological recordings of neurons</td>
<td>Decision making</td>
<td>Rat</td>
<td>Thinner neocortex did not alter decision making of the offspring</td>
</tr>
<tr>
<td>Roitman &amp; Loriaux, 2014</td>
<td>Nucleus accumbens</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>Impulsivity/Restrain (Nucleus accumbens cue responses correlated with action, regardless of cue type or accuracy)</td>
<td>—</td>
<td>Rat</td>
<td>Individual cue-responsive neurons showed either increases or decreases in activity at cue onset</td>
</tr>
<tr>
<td>Lobo et al, 2015</td>
<td>Olfactory Tubercle (Islands of Calleja, and islands of Calleja Major)</td>
<td>D&lt;sub&gt;3&lt;/sub&gt;</td>
<td>—</td>
<td>—</td>
<td>In situ hybridization (messeger ribonucleic acid levels of D&lt;sub&gt;3&lt;/sub&gt;)</td>
<td>—</td>
<td>Rat</td>
<td>messenger ribonucleic acid levels of D&lt;sub&gt;3&lt;/sub&gt; in Islands of Calleja (r = 0.91), and Islands of Calleja Major (r = 0.62) are correlated with performance in task</td>
</tr>
<tr>
<td>Ishii et al, 2012</td>
<td>Orbitofrontal cortex</td>
<td>GABA, GABA&lt;sub&gt;A&lt;/sub&gt;, and GABA&lt;sub&gt;B&lt;/sub&gt;</td>
<td>Muscimol, agonist, infusion and baclofen, agonist, infusion</td>
<td>Rodent gambling task</td>
<td>The amount of gambling task. The delay gambling task</td>
<td>Risk preference</td>
<td>Rat</td>
<td>Orbitofrontal cortex inactivation increase risk preference</td>
</tr>
<tr>
<td>Ishii et al, 2015</td>
<td>Orbitofrontal cortex</td>
<td>Serotonin, 5-HT&lt;sub&gt;1A&lt;/sub&gt;</td>
<td>WAY100635, antagonist, infusion</td>
<td>Rodent gambling task (modified)</td>
<td>—</td>
<td>Risk preference</td>
<td>Rat</td>
<td>Orbitofrontal cortex antagonism decreased risk preference</td>
</tr>
</tbody>
</table>

(Continued)
<table>
<thead>
<tr>
<th>Author &amp; year</th>
<th>Brain area</th>
<th>Messenger &amp; receptor</th>
<th>Drug name, effect, route</th>
<th>Behavioral test</th>
<th>Other technique/treatment used</th>
<th>Mental process and/or conduct analyzed</th>
<th>Species</th>
<th>Relevance of study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ishii et al, 2015</td>
<td>Orbitofrontal cortex</td>
<td>Dopamine, D&lt;sub&gt;1&lt;/sub&gt; and D&lt;sub&gt;2&lt;/sub&gt;</td>
<td>SCH 23390 hydrochloride, antagonist, infusion</td>
<td>Rodent gambling task (modified)</td>
<td>—</td>
<td>Risk preference</td>
<td>Rats</td>
<td>Orbitofrontal cortex antagonism did not alter risk preference</td>
</tr>
<tr>
<td>Ishii et al, 2015</td>
<td>Orbitofrontal cortex</td>
<td>Serotonin and 5-HT&lt;sub&gt;2A&lt;/sub&gt;</td>
<td>M100907, antagonist and infusion</td>
<td>Rodent gambling task (modified)</td>
<td>—</td>
<td>Risk preference</td>
<td>Rats</td>
<td>Orbitofrontal cortex antagonism did not alter risk preference</td>
</tr>
<tr>
<td>Zeeb et al, 2015</td>
<td>Orbitofrontal cortex</td>
<td>GABA, GABA&lt;sub&gt;A&lt;/sub&gt; and GABA&lt;sub&gt;B&lt;/sub&gt;</td>
<td>Muscimol, agonist, infusion and baclofen, agonist, infusion</td>
<td>Rodent gambling task</td>
<td>—</td>
<td>Decision making</td>
<td>Rats</td>
<td>Orbitofrontal cortex inactivation did not affect decision making</td>
</tr>
<tr>
<td>Paine et al, 2013</td>
<td>Medial prefrontal cortex</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Paine et al, 2015</td>
<td>Medial prefrontal cortex</td>
<td>GABA and GABA&lt;sub&gt;A&lt;/sub&gt;</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Baarendse et al, 2013</td>
<td>Medial prefrontal cortex</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

Disruptions in GABA dynamics contribute to decision-making deficits.

- Isolation induced lasting changes in pyramidal neurons and synapses of the medial prefrontal cortex.

(Continued)
<table>
<thead>
<tr>
<th>Author &amp; year</th>
<th>Brain area</th>
<th>Messenger &amp; receptor</th>
<th>Drug name, effect, route</th>
<th>Behavioral test</th>
<th>Other technique/treatment used</th>
<th>Mental process and/or conduct analyzed</th>
<th>Species</th>
<th>Relevance of study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fitoussi et al, 2015</td>
<td>Prefrontal – subcortical network</td>
<td>Dopamine and serotonin</td>
<td>—</td>
<td>Rodent gambling task</td>
<td>- Histology and chromatographic analyses of post-mortem sample, c-fos immuno-histochemistry</td>
<td>Decision making</td>
<td>Rats</td>
<td>Good decision: wider network, a disengagement of key prefrontal areas (insula...</td>
</tr>
<tr>
<td>Cocker et al, 2012</td>
<td>Striatum</td>
<td>Dopamine, D&lt;sub&gt;2&lt;/sub&gt; and D&lt;sub&gt;3&lt;/sub&gt;</td>
<td>—</td>
<td>Novel task for decision making</td>
<td>Micro – positron emission tomography, and autoradiography using &lt;sup&gt;11&lt;/sup&gt;C ...</td>
<td>Bet sensitivity</td>
<td>Rats</td>
<td>Low striatal D&lt;sub&gt;2&lt;/sub&gt; and D&lt;sub&gt;3&lt;/sub&gt; density correlates to high wager sensitivity</td>
</tr>
<tr>
<td>Gadziola and Wesson, 2016</td>
<td>Ventral striatum, olfactory tubercle</td>
<td>—</td>
<td>—</td>
<td>Novel water - motivated instrumental task</td>
<td>Stereotypic behavior paradigm</td>
<td>Stereotypic behavior</td>
<td>Mice</td>
<td>- Elevated activity of the ventral striatum relates to stereotypic behavior</td>
</tr>
<tr>
<td>Phillips et al, 2016</td>
<td>Ventral striatum</td>
<td>—</td>
<td>—</td>
<td>Stereotypic behavior</td>
<td>Immunohistochemical staining of FosB and ΔFosB technique</td>
<td>Stereotypic behavior</td>
<td>Mice</td>
<td>Related to the compulsivity seen in ludopathy</td>
</tr>
<tr>
<td>Aleksandrova et al, 2013</td>
<td>Subthalamic nucleus</td>
<td>—</td>
<td>—</td>
<td>Rodent gambling task</td>
<td>- Deep brain stimulation</td>
<td>Impulsivity/restrain</td>
<td>Rats</td>
<td>Sustained deep brain stimulation in the subthalamic nucleus, induced increase of premature responding in the task, and this persisted after finishing deep brain stimulation</td>
</tr>
</tbody>
</table>

**Abbreviations:** <sup>11</sup>C carbon 11; D<sub>1</sub>, dopamine 1 receptor; D<sub>2</sub>, dopamine 2 receptor; D<sub>3</sub>, dopamine 3 receptor; D<sub>4</sub>, dopamine 4 receptor; GABA, gamma-aminobutyric acid; GABA<sub>A</sub>, gamma-aminobutyric acid receptor A; GABA<sub>B</sub>, gamma-aminobutyric acid receptor B; P<sub>21</sub>, post-natal day 21; P<sub>42</sub>, post-natal day 42; 5-HT<sub>1A</sub>, serotonin 1A receptor; 5-HT<sub>2A</sub>, serotonin 2A receptor; sc, subcutaneous administration.
specie (rat or mice), author and year of publication, and relevance of the study.

Neurotransmitter

Acetylcholine receptor (cholinergic system)
The antagonism of muscarinic receptors (scopolamine, sc) but not of nicotinic receptors (mecamylamine hydrochloride, sc) impaired decision making in rat gambling tasks. Hence, muscarinic receptors can specifically disrupt decision making under conditions of risk and uncertainty (like those found in gambling).

NMDA antagonists
The blockade of NMDA receptors (but not AMPA (alpha-aminoo-3-hydroxy-5-methylisoxazolepropionate) receptors) with the antagonist MK-801 hydrogen maleate (non-competitive antagonist; sc) decreased sensitivity to delayed and uncertain reinforcement in rats, based on the delayed/probabilistic reinforcement, and on the sensitivity to reinforcer amount tests (operant conditioning chambers). Moreover, the antagonism with ketamine hydrochloride (uncompetitive antagonist, ip) decreased sensitivity to reinforcer amount without altering delay/probability discounting in the same tests. These findings suggest that NMDA receptors differentially mediate impulsivity, with MK-801 hydrogen maleate reducing impulsive choice, but augmenting risky decisions. It is relevant to consider this contrast for treating individuals displaying different psychiatric disorders characterized by impulsivity or risky decisions. In this case, a subject displaying marked impulsive choices would be treated better by means of a drug similar to MK-801 hydrogen maleate; nevertheless, the same medication might be inadequate for someone with GD.

CB₁ and CB₂
The blockade of CB₁ (agonist AM 4113, ip) or CB₂ receptors (agonist/inverse agonist AM 630, ip) or the inhibition of fatty-acid amide hydrolase (URB 597, ip) did not influence the rat gambling task performance. However, the agonism of CB₁ and CB₂ receptors (WIN 55, 212-2, ip) improved choice strategy, and increased choice latency in the suboptimal group; but only increased perseverative behavior when punished, in the optimal group. This could be interpreted as the stimulation of cannabinoid receptors could induce different gambling choice conducts based on the type of subjects; specifically, in healthy subjects (optimal group) induce inadequate conducts, but in dysfunctional subjects (suboptimal group) induced adequate conducts.

As a reference, it has been pointed out that the endocannabinoid system is associated with the reinforcing effects of drugs of abuse. CB₂ receptors have been linked to central functions, including a role in addictive processes. Moreover, CB₁ receptors are located presynaptically, inhibit synaptic transmission, and allow synaptic modulation. Furthermore, CB₂ receptors are located in different zones of the nervous system: periphery, striatum, hippocampus, thalamus, and ventral tegmental area (VTA).

DA receptors
An investigation did report that either D₄ receptors agonism (PD 168077, ip) or D₄ receptors antagonism (L-745,870, ip) had a minimal effect on latency measures and decision making, during the rodent gambling task. Additionally, neither the D₃ receptor agonism (PD 128907, ip) nor the D₃ receptor antagonism (SB 277011-A, ip) influenced decision making. Also, the antagonism of D₂ receptor (L-741,626, sc) did not affect decision making. In general, D₂, D₃, and D₄ ligands did not influence significantly the choice behaviors of the rodent gambling task.

As reference, the D₄ receptors can be found in the next areas within the nervous system: cerebral cortex, amygdala, hypothalamus, pituitary gland, visual system (retina), and the basal ganglia. Furthermore, D₃ receptors are localized in the islands of Calleja, mammillary bodies, nucleus accumbens (NAc) shell, frontoparietal cortex, the substantia nigra/VTA, basolateral amygdala (BLA), and lateral habenula. In general, some authors agree that the specific localizations of D₂ and D₄ dopamine receptors in the nervous system support their roles in cognition and emotion.

Another investigation reported that pramipexole (an agonist of D₂ and D₃ receptors, sc) induced GD tendencies based on a probability discounting task in rats. Specifically, pramipexole augmented unfavorable decisions, disrupted the discounting of probabilistic losses, augmented risk-taking behaviors, distorted the representation of rewards, and impaired the ability to discern favorable from unfavorable contingencies. Moreover, the results of complementary studies (voltammetry recordings and High Performance Liquid Chromatography (HPLC)) focused in the NAc suggested that pramipexole behavioral effects were separated from the dynamic changes related to mesolimbic DA release.
As a reference, the HPLC also measured besides dopamine level, the level of serotonin and norepinephrine. Moreover, the pump speed (Shimadzu LC-6A liquid chromatograph, Columbia, Maryland, United States of America) was 1.5 mL/min. The reverse-phase column utilized was a Rexchrm (Regis Technologies, Morton Grove, Illinois, United States of America) S50100-ODS C18 column with a length of 25 cm and an internal diameter of 4.6 mm. The compounds were measured at +0.7 V using a Shimadzu L-ECD-6A electrochemical detector.

Brain structures

Amygdala

Rat studies did show that amygdala low serotonergic metabolism, or its sustained activity related to poor decision making in the rat gambling task. Moreover, the lesions of the rat BLA related to reduced risk seeking for losses, but intact risk aversion for gains, based on the loss – chasing task, and the betting task; these data supported the hypothesis that the amygdala plays a more prominent role in choice biases related to losses. The Tremblay’s data suggested that risk seeking for losses being explained by changes in the amygdalar activity, because the amygdala roles in representing the negative affect, and the aversive emotional reaction to loss. Also, these findings discouraged the explanation of risk seeking for losses, because the aberrant estimations of probability or loss magnitude. This result suggested that communication between these areas is vital for the appropriate assessment of reward value to influence choice.

In addition, another rat investigation explored the disconnection between BLA pathways and the OFC, and found a retarded acquisition in the gambling task. Based on Zeeb opinion, this disconnection prevented modifications in the value of a specific reward for contributing appropriately to cost-benefit decision making. Also, it seems that pathways from the OFC to BLA are important in the decision process, and for the adequate assessment of reward value to influence choice.

As a reference, a rat study reported that this species has decision-making processes that are influenced by a previous reference point; this study used a modified T maze paradigm. Specifically, the modification did consist of adding “pockets” at both sides (right and left) of the T’s stem; because, these pockets in the stem stored pellets, rats could set reference values for each arm of the maze, before selecting.

The “previous expectation” had been previously reported in human research. It is known that decision-making processes can be disrupted in GD. If the decision making can be influenced by the “previous expectation”, then this expectation should be studied, and its neurobiology for easing the treatment of GD. It is still necessary to find out which brain structure(s) is(are) involved in a “previous expectation”.

Cingulate cortex

Researches regarding the inactivation of the anterior cingulate cortex (ACC) by means of GABA_A and GABA_B agonism reported opposed results. Specifically, one study stated that inactivation of the ACC by means of a mixture of the GABA_A agonist (muscimol; infusion in the brain) and the GABA_B agonist (baclofen; infusion in the brain) impaired rodent’s ability to differentiate winning from losing outcomes in a rat slot machine task. However, the other study that inactivated ACC by means of GABA_B (baclofen hydrochloride, brain microinfusion) and GABA_A (muscimol hydrobromide, brain microinfusion) receptors agonism reported no effect on decision making based on a rat gambling task.

Moreover, another study explored the effect of D_4 agonism (PD168077; infusion in the brain) in the ACC, and it found a disruption of the rat’s ability to differentiate winning from losing outcomes in the slot machine task. Also, it found an augmentation of the reward expectancy, but only on archetypal “near-miss” trials (ie, when the first two of three stimuli in the array were concordant with a rewarding outcome, and only the last stimulus critically signaled a non-win); Cocker considered that the ACC is fundamental for analyzing the adequate response when competing stimulus and outcome associations are activated; also, this author considered that the D_4 receptor antagonists might be an effective treatment for GD.

Hippocampus

High levels of DA, 5-HT, and noradrenaline in the hippocampus predicted the emergence of more exploratory and risky behaviors in a strain of healthy inbred mice, based on a gambling task. In this study, they focused on postmortem brain analysis, rather than administration of drugs, or experimental treatments on living brains.

Insular cortex

Some studies evaluated different areas of the insular cortex like: overall insular cortex, anterior insular cortex, or agranular insular cortex (rostral or caudal). The insular cortex
did seem relevant to the rapid (30 mins) disruptive action of corticosteroid hormones (C174 Corticosterone HBC-complex, sc) on decisions (reward based) of the rat’s Iowa gambling task.\textsuperscript{51} This corticosteroid action was related to stress experience. As a reference, the disruption on decision was accompanied by significant changes in the insular cortex (based on c-fos immuno-histochemistry).

Another study showed that inactivation of insular cortex by a mixture of GABA\textsubscript{A} (muscimol; brain microinjection) and GABA\textsubscript{B} (baclofen hydrochloride; brain microinjection) receptors agonists induced risky behaviors linked to altered decisions, based on a rat gambling test (a radial arm maze).\textsuperscript{52}

Other studies investigated the anterior insular cortex relevance in gambling-related behaviors, and found mixed results. Specifically, the first investigation found a decreased in risk preference, based on two rats gambling tasks (the amount gambling task and the delay gambling task); the treatment was a mixture of GABA\textsubscript{A} (muscimol; brain microinjection) and GABA\textsubscript{B} (baclofen; brain microinjection) receptor agonists.\textsuperscript{53} Moreover, the second investigation blocked the D\textsubscript{2} receptors of the anterior insular cortex (etaclopride hydrochloride, brain microinjection) and found augmentation of risk preference, after winning in a previous risky choice; also, the blockade of the 5-HT\textsubscript{1A} receptors (WAY100635; brain microinjection) of the anterior insular cortex increased risk preference, after losing in a previous risky choice of a gambling task.\textsuperscript{54} As reference, the antagonism of dopamine 1 receptor (D\textsubscript{1}) (SCH23390 hydrochloride) or the antagonism of serotonin 2A receptor (5-HT\textsubscript{2A}) (M100907; brain microinjection) in the anterior insular cortex did not alter risk preference in the rat gambling task.\textsuperscript{54}

Regarding the agranular insular cortex, its inactivation by a mixture of GABA\textsubscript{A} (muscimol, brain microinjection) and GABA\textsubscript{B} receptors (baclofen, brain microinjection) agonism decrease risk preference based on two different rat gambling tasks; besides, in risk-free control situations, the agranular insular cortex inactivation did not impair decision making.\textsuperscript{53} Furthermore, the inactivation of the caudal agranular insular cortex by either lesion (ibotenic acid; brain microinjection) or by a mixture of GABA\textsubscript{A} (muscimol; brain microinjection) and GABA\textsubscript{B} (baclofen; brain microinjection) receptors agonists did not disrupt decision-making behavior under risk in a rat gambling task.\textsuperscript{55}

**Lateral ventricles**

Brain anatomical abnormalities like the enlargement of the lateral ventricles did not alter decision making in the rat gambling task.\textsuperscript{56} As a reference, the enlargement of ventricles was induced by a maternal (before and during pregnancy) diet deficient in vitamin D; subsequently, the whole litters were placed in a standard diet and evaluated.

**Limbic area**

Different experimental manipulations of the infralimbic area generated poor decision making in the rat gambling task; for instance: a higher serotonergic metabolism, a rapid action of corticosteroids hormones (30 mins), and inactivation by gamma-aminobutyric acid (GABA) receptors agonism. Specifically, the relationships between a higher serotonergic metabolism in the infralimbic area, and poor decision making in the rat gambling task was inferred based on the postmortem brain analysis after gambling behavioral tests.\textsuperscript{44} Moreover, the rapid and disruptive action of corticosteroid hormones of stress (C174 Corticosterone HBC-complex; sc) on decision making of rats was performed by means of non-invasive brain manipulations.\textsuperscript{51} Furthermore, the disruption of decision making after agonism of GABA\textsubscript{A} (muscimol hydrobromide; brain microinjection) and GABA\textsubscript{B} (baclofen hydrochloride; brain microinjection) receptors was performed by means of direct brain injections; specifically, this inactivation allowed an augmented preference for disadvantageous options and reduced choice for optimal options.\textsuperscript{49}

However, other conditions like manipulations of the D\textsubscript{2} receptors did not alter neither choice preference nor optimal performance in the rat gambling task; specifically, the infralimbic area generated poor decision making by means of administering the D\textsubscript{2} receptor antagonist (etaclopride hydrochloride; brain microinjection).\textsuperscript{49}

On the other hand, another set of studies targeted the prelimbic cortex by the agonism of GABAergic receptors and the antagonism of D\textsubscript{2} receptors obtaining opposed results.\textsuperscript{49} Specifically, the inactivation of prelimbic cortex by means of the agonism of GABA\textsubscript{A} (muscimol hydrobromide; brain microinjection) and GABA\textsubscript{B} (baclofen hydrochloride; brain microinjection) receptors disrupted decision making in the rat gambling task.\textsuperscript{49} Moreover, this inactivation allowed an augmented choice for disadvantageous options, and a reduced choice for optimal options.\textsuperscript{49} Under other conditions, the disruption of the prelimbic cortex by means of treatment with the D\textsubscript{2} receptor antagonist (etaclopride hydrochloride; brain microinjection) did not alter choice preference neither optimal performance in the rat gambling task.\textsuperscript{49}
Neocortex
Nervous system anatomical abnormalities like a tiny cerebral cortex did not modify decision making in the rat gambling task. As a reference, the reduction of the cerebral cortex was produced by a maternal diet (before and during pregnancy) deficient in vitamin D; afterward, the offsprings were placed in a standard diet and evaluated.

NAc
A study found a relationship between the activity of specific cue responsive NAc neurons, and the cue onset during a go/no go task. Despite a gambling task was not used, the go/no go task has relevance to impulsivity, that is a core trait of GD. Specifically, electrophysiological recordings of neurons in the NAc during the go/no go tasks showed that individual cue-responsive neurons displayed either increases or decreases in activity at the cue onset; NAc cue responses correlated with action, regardless of cue type or accuracy.

Olfactory tubercle
An investigation found a correlation between the messenger ribonucleic acid (mRNA) levels of D3 receptors in the island of Calleja (r=−0.91), in the islands of Calleja major (r=0.62) and the performance of male rats in the rodent gambling task (this study only used males). This finding was consistent with a human imaging study (positron emission tomography) that reported a link between D3 receptor binding index and the severity of disordered gambling.

OFC
A group of studies with different manipulations (neurotransmitters and receptors systems) reported diverse results: eg, increase in risk preference (inactivation by GABAergic agonism, or rapid corticosteroid action), decrease in risk preference (antagonism of 5-HT1A), no effects on risk preference (antagonism of D1, D2, or 5-HT2A receptors), and no effect on decision making (inactivation by GABAergic agonism or D2 receptor antagonism). However, it is important to note that under risk-free control situations, the GABAergic agonism of OFC did not affect decision making; hence, the degree of risk of the task should be considered. Specific details of all the previous reports are explained in the next paragraphs.

The OFC was inactivated by a mixture of GABA_A (muscimol; brain microinjection) and GABA_B (baclofen; brain microinjection) receptors agonists; this treatment augmented risk preference in the rats, based on two gambling tasks (the amount gambling task, and the delay gambling task). However, under risk-free control situations, the inhibition of the OFC did not disrupt decision making. According to the authors, the OFC denoted relevance at the time of accepting or declining a risk. Moreover, the lateral OFC did show relevance for the rapid (30 mins) disruptive action of corticosteroid hormone (C174 Corticosterone HBC-complex; sc) on decision making (reward based) in a rat Iowa gambling task. This was inferred because the disruption on decision process was accompanied by significant changes in gene expression in the lateral OFC (increase in c-fos expression, based on c-fos immuno-histochemistry).

Nevertheless, the antagonism of 5-HT1A receptors (WAY100635; brain microinjection) in the rat’s OFC decreased risk preference on a modified gambling task. Finally, other studies reported absence of effects of OFC manipulation; specifically, the antagonism of either D1 receptors (SCH 23390 hydrochloride; brain microinjection), D2 receptors (eticlopride hydrochloride; brain microinjection), or 5-HT2A receptors (M100907; brain microinjection) did not alter risk preference on a modified gambling task. Furthermore, the inactivation of the OFC by means of GABA_A (muscimol hydrobromide; brain microinjection) and GABA_B (baclofen hydrochloride; brain microinjection) receptors did not affect decision making in rats, based on a gambling task. Moreover, D2 receptor antagonism (eticlopride hydrochloride; brain microinjection) did not affect decision making in the same paradigm.

mPFC
Some studies showed that direct manipulation (ibotenic acid lesion, GABAergic antagonism) or developmental manipulation (adolescence/juvenile social isolation) of the mPFC disrupted decision making. More details about the previous reports are explained in the next paragraphs.

First, the excitotoxic lesion of mPFC induced by ibotenic acid (brain microinjection) worsened decision making (fewer selection of advantageous or optimal choices) based on a rat gambling task; however, this deficit in decision making was attenuated after treatment with D1 receptors antagonism (SCH23390; ip). However, the D2 receptors antagonism (haloperidol; sc) did not attenuate the deficit.
Furthermore, the antagonism of GABA_{A} receptors (bicuculline methiodide; brain microinjection) in the mPFC disrupted decision making in the rat gambling task. Despite this study described this application for schizophrenia treatment, this finding is also useful for GD treatment, because it studied decision process during the rat gambling task.61 Finally, social isolation from early adolescent to juvenile period (post-natal day 21 (P_{21}) to post-natal day 42 (P_{42})) induced lasting cellular and synaptic changes in the pyramidal neurons of the adult mPFC.62 Besides, isolation consequences counteract the DA enhancement induced by a DA agonism bolsterer (amphetamine sulfate; ip) or by a DA reuptake inhibitor (GBR12909 dihydrochloride; ip) in the five-choice serial reaction time task (challenging conditions).62 Also, the social isolation decreased sensitivity to DA in the pyramidal neurons of the mPFC. Impulsivity was measured in the rat gambling task and other tests. Also, social isolation impaired impulsive action and decision making under novel or challenging circumstances based on the rat gambling task and other tests. However, impulsive choices were not affected by social isolation.62

PFC – subcortical network and related structures
A rats’ study combined gambling tasks, post-mortem analysis (DA and 5-HT turnovers), and c-fos immuno-detection in the brain prefrontal – subcortical network.44 Differences between good and bad decision making was found. Good decision making was characterized by a wider network (but once good choices had been made), and a disengagement of the key prefrontal areas (insula and infralimbic cortices) and the amygdala. On the other hand, poor decision making was related to a lower network recruitment and to a sustained amygdala activity.44 Besides, poor decision making was linked to an imbalance of monoaminergic metabolism (ie: a higher infralimbic vs a lower amygdalar serotonergic metabolism), and to an aberrant low recruitment of brain areas linked to executive functions and affective valence during decision processes.44

Striatum
Studies have looked at the relevance of the striatum in general (rats), and its specific zones like olfactory tubercle and ventral striatum (mice). In general, it was found that striatum activity was linked to wager sensitivity, motivated behavior, discrimination of rewards, stereotypical behavior, and compulsivity.53–65 Additional technical details are described in the next paragraphs.

Specifically, lower striatal D_{2} and D_{3} receptors densities correlated to high wager sensitivity, based on a novel task for decision making in rats, micro-positron emission tomography, and autoradiography using [11C] raclopride.63

In addition, a mice study reported that the olfactory tubercle (a sub-region of the ventral striatum) robustly encoded the onset and progression of motivated behaviors (organization of goal-directed behaviors), and discriminated the type and magnitude of a reward (process of reward information).64 As reference, this mice investigation did use a novel water-motivated instrumental task, and “in vivo” electrophysiological recordings; despite this investigation did not perform explicit behavioral tests about gambling, it was proposed by the authors, that the findings were conceptually/theoretically related to GD.64 Finally, another investigation performed on mice found that an augmented activity of the ventral striatum related to stereotypical behavior.65 The mice were evaluated by means of a stereotypical behavior paradigm, and the brains analyzed by immunohistochemical staining of FosB and delta FosB.65 The authors (Phillips et al) proposed that the stereotypy observed could be relevant to the compulsivity described in GD and other disorders (eating and drug seeking).65

STN
There was scarce research regarding the involvement of the STN in gambling behavior. Specifically, a study reported that several sessions of bilateral deep brain stimulation (DBS) of the STN induced a subsequent increment of premature responding in the gambling task; this increment even persisted after finishing the stimulation.66 As a reference, DBS of the STN had been also associated with impulsivity in the absence of Parkinsonism (under specific conditions).66

Discussion
The main question of this review was: which are the main neurotransmitters systems and brain structures relevant for GD based on recent rodent studies? This question was answered in this section by contrasting the present review main points and those from previous literature reviews in the field (reviews cited in the Introduction). The present review found that NMDA receptor antagonism influence reinforcement sensitivity and impulsivity; also, that D_{2} and D_{3} receptors’ agonism induces GD tendencies. These points agree with the publications review by Grant et al.2 it concluded that probably diverse neural systems
participate in the pathophysiology of GD like those related to glutamate and DA among others messengers.\textsuperscript{2}

The present work considers that the BLA–OFC pathway is relevant for the assessment of reward among other functions in the rat gambling task. This partially agrees with the Levy and Glimcher’s review;\textsuperscript{13} precisely, those authors proposed that the vmPFC/OFC is part of a brain network that codify the values of rewards by means of a standard neural scale (based on human neuroimaging).\textsuperscript{13} It seems that the OFC is related to the assessment and codification of rewards in gambling activities.

Based on the present review, a higher level of DA, noradrenaline, and 5-HT in the hippocampus predicted exploratory and risky behaviors in gambling. Related to this, a review by Meng et al pointed out that bilateral overactivity of the parahippocampus among other structures, positively correlated with South Oaks Gambling Screen scores.\textsuperscript{14} Taking these together, it seems that a higher metabolism and activity of the zone of the hippocampus and its surroundings (parahippocampus) relates to more risky gambling tendencies.

Furthermore, the present review found that insular cortex activity relates to decision making in the rat gambling task; in the same sense, a previous literature review stated that the insular cortex (including rostral agranular zone) among difference structures influence the correctness of gambling decision.\textsuperscript{6} Taking these together, it seems that insular cortex relates to decision making in gambling tasks.

Moreover, the present review found that infralimbic area relates to decision making in the rat gambling task; this agrees with another publications review that states that infralimbic area among other structures is involved in the correctness of gambling decision on rodent tasks.\textsuperscript{6}

The present review found that OFC activity is related to risk preference; relevant to this, another review states that alterations in the OFC among other structures are found in fellows with GD.\textsuperscript{6} It seems that alteration of the OFC activity is relevant for GD. Additionally, this review found that mPFC is involved in decision making, and its disruption impairs decision making. Other reviews have proposed similar ideas; Goulet-Kennedy et al’s review pointed that PFC among other structures is fundamental for decision processes based on clinical studies.\textsuperscript{11} Moreover, a review by Potenza states that vmPFC among other areas is relevant for GD.\textsuperscript{15} In general, it seems that PFC (including medial and ventromedial area) is relevant in the dynamics of GD.

Besides, the present review points that PFC–subcortex network activity is linked to poor decision making if lower network action and sustained activity of the amygdala are present; moreover, PFC–subcortex network is associated with good decision if a wider network and disengagement of key prefrontal areas and the amygdala are present. These points agree with Grant et al’s review;\textsuperscript{2} Grant et al work concludes that based on imaging reports, GD relates to anatomical and functional anomalies of nexus involved in reward processing and top-down monitoring.\textsuperscript{2} Hence, both reviews agree that disruption of top-down circuits is a common element in problems linked to gambling.

Regarding the striatum, the present literature review states that the striatum’s density of dopamine receptors relates to wager sensitivity; also, the ventral striatum activity relates to stereotypy (like the GD compulsivity). Moreover, the olfactory tubercle relates to the onset and progression of motivated behaviors and reward’s discrimination. Similarly, other publication reviews like the one by Goulet-Kennedy et al pointed that the striatum is a conductor of decision processes (which are relevant to gambling behaviors) based on clinical studies.\textsuperscript{11}

Furthermore, the review by Norbury and Husain states that marked sensation seeking relates to GD, and to dopaminergic transmission;\textsuperscript{5} specifically, fellows with marked sensation seeking display high tonic DA levels and over-responsive midbrain dopaminergic responses to signals of future reward.\textsuperscript{5} Also, differences in subject reactions (variability in approach – avoidance reactions) to stimuli stems from differences in the efficiency of DA transmission at the level of striatum.\textsuperscript{5} Another review by Potenza, stresses that the ventral striatum among other structures is linked to gambling and GD.\textsuperscript{15} Integrating, the reviews state that striatum relates to wager sensitivity (based on DA receptor density), stereotypy (ventral zone), conduction of decision processes (including those of gambling behaviors), and variability in approach – avoidance to stimuli.

The different points contrasted above in the Discussion section, between the present and other reviews published, have been integrated for elaborating clinical indications. These indications are the next: a) glutamate and DA seem relevant in the pathophysiology of GD; however, other neurotransmitters should also be considered, b) the OFC is relevant for the assessment and codification of rewards in gambling activities, c) a higher metabolism and activity of the hippocampus and its surroundings (parahippocampus) relates to risky gambling tendencies, d) the insular cortex and the infralimbic area are relevant for gambling-
related decisions, e) the alteration of OFC activity is relevant for GD, f) the PFC (including mPFC and vmPFC) is relevant for the dynamic of GD, g) the disruption of top (cortical)–down (subcortical) circuits can be relevant to gambling problems, and h) the striatum relates to wager sensitivity, stereotypy, decision processes, and approach/avoidance to stimuli related to gambling.

**Conclusion**

Based on the studies revised that used noninvasive methods for drug administration, some of the receptors involved in behaviors related to GD are: muscarinic, NMDA, CB1, CB2, D1, D3, and D4 receptors. Moreover, based on the studies revised that used invasive methods for drug administration, some of the neurotransmitters and receptors involved in GD are: 5-HT1A, noradrenaline receptors, GABA A, and GABA B. According to this work, the next brain structures are involved in behaviors related to GD: amygdala (including BLA), BLA–OFC pathways, ACC, hippocampus, infralimbic area, prelimbic cortex, insular cortex (including anterior and rostral agranular zones), NAc, olfactory tubercle (island of Calleja and the island of Calleja major), OFC, mPFC, PFC–subcortical network, striatum (including ventral zone and olfactory tubercle), and STN. The present review and others described in the field agree that DA and glutamate, among other neurotransmitters, are relevant to GD. The present review and others described in the field agree that the next brain areas are relevant for GD: OFC, hippocampus/parahippocampus, insular cortex, infralimbic area, PFC, PFC–subcortical network, and striatum. The search for GD treatments should consider and integrate this diversity of neurotransmitters, receptors, and brain areas.

**Abbreviation list**

AMPA, alpha-amino-3-hydroxy-5-methylisoxazolepropionate; ACC, anterior cingulate cortex; BLA, basolateral amygdala; CB1, cannabinoid receptor 1; CB2, cannabinoid receptor 2; DBS, deep brain stimulation; DSM-5, Diagnostic and Statistical Manual of Mental Disorders-5; DA, dopamine; D1, dopamine 1 receptor; D2, dopamine 2 receptor; D3, dopamine 3 receptor; D4, dopamine 4 receptor; EEG, electroencephalography; GD, gambling disorder; GABA, gamma-aminobutyric acid; GABA A, gamma-aminobutyric acid receptor A; GABA B, gamma-aminobutyric acid receptor B; HPLC, High Performance Liquid Chromatography; ip, intraperitoneal administration; mPFC, medial prefrontal cortex; mRNA, messenger ribonucleic acid; mins, minutes; NMDA, N-methyl-D-aspartate; NAc, nucleus accumbens; OFC, orbitofrontal cortex; P21, post-natal day 21; P42, post-natal day 42; PFC, prefrontal cortex; SENACYT, Secretaria Nacional de Ciencia, Tecnologia e Innovacion (English: National Secretariat of Science, Technology and Innovation); 5-HT, serotonin; 5-HT1A, serotonin 1A receptor; 5-HT2A, serotonin 2A receptor; sc, subcutaneous administration; SNI, Sistema Nacional de Investigacion (English: National System of Investigation); STN, subthalamic nucleus; VTA, ventral tegmental area; vmPFC, ventromedial prefrontal cortex.

**Acknowledgments**

We would like to thank Open Access Publishing Fund from Florida State University Libraries for its funding. Dr. G. Quintero Garzola belongs and is backed by the SNI (Sistema Nacional de Investigacion) from SENACYT. SENACYT is a public organization located in Panama.

**Disclosure**

The author reports no conflicts of interest in this work.

**References**

Dovepress

Quintero Garzola

Neuropsychiatric Disease and Treatment 2019:15

neuropharmacol.2016.11.014


Dovepress

Neuropsychiatric Disease and Treatment downloaded from https://www.dovepress.com/ by 54.70.40.11 on 10-Nov-2019

For personal use only.