Review

The structural and functional connectivity of the amygdala: From normal emotion to pathological anxiety

M. Justin Kim*, Rebecca A. Loucks, Amy L. Palmer, Annemarie C. Brown, Kimberly M. Solomon, Ashley N. Marchante, Paul J. Whalen

Department of Psychological & Brain Sciences, Dartmouth College, 6207 Moore Hall, Hanover, NH 03755, USA

A R T I C L E   I N F O

Article history:
Received 3 April 2011
Accepted 16 April 2011
Available online 22 April 2011

Keywords:
Amygdala
Medial prefrontal cortex
Connectivity
Anxiety
Emotion disorders
Emotion regulation

A B S T R A C T

The dynamic interactions between the amygdala and the medial prefrontal cortex (mPFC) are usefully conceptualized as a circuit that both allows us to react automatically to biologically relevant predictive stimuli as well as regulate these reactions when the situation calls for it. In this review, we will begin by discussing the role of this amygdala–mPFC circuitry in the conditioning and extinction of aversive learning in animals. We will then relate these data to emotional regulation paradigms in humans. Finally, we will consider how these processes are compromised in normal and pathological anxiety. We conclude that the capacity for efficient crosstalk between the amygdala and the mPFC, which is represented as the strength of the amygdala–mPFC circuitry, is crucial to beneficial outcomes in terms of reported anxiety.

© 2011 Elsevier B.V. All rights reserved.

Contents

1. Introduction .................................................................................................................. 403
2. The structural and functional connectivity of the human amygdala and prefrontal cortex ........................................................................................................... 404
  2.1. Structural neuroanatomy of amygdala–mPFC circuitry ........................................... 404
  2.2. Functional neuroanatomy of the amygdala–mPFC circuitry ........................................ 404
  2.3. Amygdala–mPFC circuitry at rest ........................................................................... 405
3. Amygdala–prefrontal circuitry and fear conditioning and extinction .............................. 405
4. Amygdala–prefrontal circuitry and emotion regulation .................................................. 405
5. Amygdala–prefrontal circuitry and the interpretation of emotionally ambiguous facial expressions .......................................................... 406
6. Amygdala–prefrontal circuitry and anxiety within the normal range ............................... 407
7. Amygdala–prefrontal circuitry and pathological anxiety ............................................. 407
  7.1. Social anxiety disorder ...................................................................................... 407
  7.2. Posttraumatic stress disorder ........................................................................... 408
8. Conclusions ................................................................................................................. 408
Acknowledgements ......................................................................................................... 408
References ........................................................................................................................ 408

1. Introduction

Accurate evaluation of and response to potentially life-threatening or sustaining events are hallmarks of biologically relevant learning in animals and humans. In response to cues of threat, rodents exhibit a distinctive “freezing” or somatomotor arrest behavior. This behavior is critical in a natural environment in which movement may attract a predator to its location. Humans show a similar freezing response to a potentially threatening situation [1]. Rather than having to avoid predators, humans might more ordinarily show such a response when having to speak in front of a large audience. In such threatening instances, performance would be facilitated if able to override the initial freezing behavior.

In psychological terms, instinctive reactions to threat and subsequent regulatory responses are often referred to as bottom-up and top-down processes, respectively. The interplay between these two processes is exemplified by the following example: upon encountering a snake at a zoo, an initial reaction is driven by its appearance (i.e., bottom-up saliency), but the response is then implicitly con-
trolled by the determination that the snake presents no immediate danger because it is behind a sheet of Plexiglas (i.e., top-down control). Of course, the context is critical since the same snake encountered in a field would evoke an initial freezing response followed by a very different type of top-down control in the form of running (or screaming in some cases). Thus, interactions between bottom-up and top-down processes will determine the adaptiveness of behavior in a given situation.

This conceptualization may be directly applicable to clinical research, as the interaction between these bottom-up and top-down processes is hypothesized to be impaired in psychiatric illnesses – and here we will focus on the anxiety disorders. For example, in specific phobias, perhaps a failure to employ top-down control mechanisms allows initial bottom-up responses to intrude on normal cognitive functioning. Alternatively, it may be the case that the initial bottom-up reactions are so potent and exaggerated that even a normally functioning top-down regulatory system cannot keep these responses in check. Individual differences in the function and structure of this circuitry can also explain differences in normal levels of anxiety.

Numerous studies have highlighted the critical role of the amygdala and the mPFC in behavioral phenomena that involve competition between bottom-up and top-down processes, including fear conditioning and extinction [2–4]. Critically, it is believed that the mPFC regulates and controls amygdala output and the accompanying behavioral phenomena [2–4]. The reciprocal relationship between the amygdala and the mPFC strongly suggests the need to investigate these brain regions as one circuit, rather than studying them separately. That is, while numerous studies have assessed the separate contributions that the amygdala and mPFC make to reactivity and regulation, respectively [5–8], more recent studies suggest that the structural and functional connectivity between these two regions is a better predictor of these outcomes than the activity of either region alone [9–11]. The idea here is that the stronger the coupling between the amygdala and the mPFC, the better the behavioral outcome in terms of reported anxiety.

2. The structural and functional connectivity of the human amygdala and prefrontal cortex

2.1. Structural neuroanatomy of amygdala–mPFC circuitry

The amygdala is an almond-shaped brain structure that resides in the medial temporal lobe of the brain [12,13]. Its structure is comprised of many subnuclei, including the basolateral nuclei (BLA) and the central nucleus (Ce), which have distinct anatomical connections with other brain regions that serve different functions. Comprehensive descriptions of the anatomical connections of the amygdala exist elsewhere [14,15]. Here, we focus on the connectivity between the amygdala and the prefrontal cortex, especially the mPFC. The mPFC can be roughly divided into two subregions, relative to the genu of the corpus callosum–dorsal mPFC (dmPFC) and the ventral mPFC (vmPFC). Broadly defined, the dmPFC includes the supragenual anterior cingulate and the medial frontal gyrus, whereas the vmPFC includes the subgenual anterior cingulate, ventromedial prefrontal and medial orbitofrontal cortex (Fig. 1).

Most of the known facts about the anatomical connections of the amygdala–mPFC circuitry are derived from animal studies, especially non-human primates. This is because the invasive nature of the methods that are used to investigate brain connections – such as lesion and tracing studies – are difficult to employ in humans. Data from non-human primate brains show that the majority of the afferent fibers to the amygdala originate in the orbitofrontal cortex and the mPFC, and these projections are denser and heavier from the caudal compared to the rostral aspects of these prefrontal areas [16–21]. In turn, the amygdala sends efferent projections to these orbitofrontal and mPFC regions [17–19,22,23], and interestingly these projections are heavier than the reciprocal cortical afferents [19,24]. Most of the amygdala–prefrontal connections are concentrated in the BLA, as opposed to Ce. Based on animal studies of fear conditioning and extinction, mPFC input to the BLA as well as the intercalated cells (adjacent to the BLA) is responsible for inhibiting amygdala output by regulating BLA inputs to the Ce (see Section 3, for details).

In humans, the structural connections of the amygdala have been investigated more recently using non-invasive neuroimaging methods such as diffusion tensor imaging (DTI). This imaging method takes advantage of the fact that the movement of water molecules in the brain differs in different types of brain tissue. To elaborate, in an unrestricted tissue environment, such as in the ventricles of the brain, water molecules show isotropic diffusion in all directions equally. Importantly, however, the movement of water molecules is greatly restricted in myelinated axons – that is, water molecules in white matter tend to move in a single direction along the myelinated axons. Thus DTI is a method optimized to assess white matter fiber tracts in the brain. There are two different types of information about which DTI can inform us – (1) the orientation of white matter fiber tracts, and (2) the strength or integrity of white matter fiber tracts. The former can be accomplished by using fiber tracking or tractography to measure the direction of water diffusion [25]. The latter can be calculated by measuring the degree of anisotropic diffusion [26]. Specifically, normalized measures such as fractional anisotropy can be computed for each brain voxel and used to index the structural integrity of the measured white matter fiber tracts [26]. A number of studies have utilized these methods to identify an amygdala–prefrontal pathway in the human with a specific focus on connectivity with the dorsal and ventral aspects of the mPFC [27–29].

2.2. Functional neuroanatomy of the amygdala–mPFC circuitry

The functional counterpart of structural connectivity – functional brain connectivity – can also be used to investigate
amygdala–prefrontal interactions. This method is optimal for understanding the relationship between spatially remote brain regions by assessing brain activity across time. Analyses of functional brain connectivity can be defined in two ways: (1) functional connectivity and (2) effective connectivity [30]. Functional connectivity is simply a measure of the temporal correlation of brain activity in two or more regions, whereas effective connectivity seeks to reveal the directional effect of one neuronal system exerted over another [30]. By definition, functional connectivity is purely correlational in nature, and provides no information regarding the directionality of how one brain region affects another. In contrast, effective connectivity attempts to explain the causal relationship between the interactions of different brain regions, relying on more advanced statistical modeling methods such as structural equation modeling [31], psychophysiological interactions [32], and dynamic causal modeling [33]. Based on the extensive anatomical connections between the amygdala and mPFC shown in human and non-human primates, a number of investigations have used these functional and effective connectivity measures to assess the strength of amygdala–mPFC coupling and its relationship with behavioral outcomes [9,34,35].

3. Amygdala-prefrontal circuitry and fear conditioning and extinction

Studies of the non-human animal amygdala have shown that sensory information received by the BLA is then passed to the Ce [45]. Though outputs exist at the level of the BLA, a majority of outputs originate from the Ce [46]. The Ce projects directly to the hypothalamus and brain stem nuclei that drive autonomic and somatomotor responding [47]. The Ce also projects to all major neuromodulatory systems including dopaminergic, cholinergic, serotonergic and noradrenergic systems [46]. Thus, while direct projections can primarily affect physiological and motor responses, these neuromodulatory projections can serve to globally, non-specifically and instantaneously effect neuronal excitability across the brain. Such changes could serve to induce a state of heightened vigilance rendering the organism a more efficient consumer of information in biologically relevant learning situations [46]. One such situation involves the acquisition and expression of learned responses through classical conditioning [48,49]. For example, in a typical aversive conditioning paradigm, subjects learn that a previously neutral stimulus (e.g., tone) predicts the occurrence of an unconditioned stimulus (US; e.g., electric shock), thereby acquiring the value of a conditioned stimulus (CS) which now elicits a conditioned response (CR; e.g., freezing) to the CS that was previously reserved for the US [50,51]. The generation of these CS–US associations and their behavioral expressions are known to be amygdala-dependent since manipulations of this structure block or retard such learning [52,53].

A reversal of this classical conditioning procedure is known as extinction – suppressing previously learned CS–US associations [54,55]. The inhibition of CS–US associations can be achieved by top-down regulatory input from the mPFC to the BLA [56]. This process is supported by the existence of amygdala–mPFC connectivity that allows direct, reciprocal communication [14,19,56]. For example, electric stimulation of the mPFC resulted in the inhibition conditioned responses, emulating the effects of extinction in the rat [56]. In humans, greater cortical thickness of the vmPFC was associated with better behavioral performance during extinction recall [57,58]. Similar findings have also been demonstrated in humans using fMRI, highlighted by increased vmPFC activity during successful extinction of learned US-CS associations [59–61]. Interestingly, a study using functional connectivity methods [59] showed that the amygdala and the vmPFC were functionally coupled during the entire course of the experiment, which included a combination of fear extinction and emotion regulation tasks. Thus, these structural and functional findings highlight the importance of amygdala–mPFC interactions for the regulation and inhibition necessary for extinction learning and/or memory.

4. Amygdala-prefrontal circuitry and emotion regulation

The ability to regulate our emotions is essential in our everyday lives, and successful emotion regulation begets beneficial outcomes in many social situations. Emotion regulation is a classic example of how top-down and bottom-up processes compete and interact to produce optimal (or counterproductive) behavioral outcomes. For example, one’s instinctive reaction to a frightening scene in a horror movie may include an urge to scream and/or run out of the room. Normally, this bottom-up reaction is controlled by a top-down intervention (e.g., reminding oneself that this is only a movie). Taking the scenario described above into account, it would not be too difficult to imagine that individuals may employ different strategies to achieve such emotion regulation.

To date, studies investigating the neural basis of emotion regulation have primarily examined two distinctive means of
emotion regulation – by simply suppressing what one is feeling (i.e., suppression), or by cognitively reevaluating the stimulus that is evoking the emotion (i.e., reappraisal) [3]. Not surprisingly, emotion regulation and the extinction of fear conditioning are suggested to have overlapping underlying neural mechanisms, since the essence of both processes involves reevaluating biologically relevant stimuli [4,62]. Like extinction, it is useful to assume that during emotional regulation the prefrontal cortex exerts control over the amygdala in response to an emotional challenge [3,63]. Based on this framework, numerous functional neuroimaging studies have demonstrated increased prefrontal activity and concomitant decreased amygdala activity during successful emotion regulation [59,64–71]. Unlike extinction, many emotion regulation studies point to the ventral and dorsal lateral prefrontal cortex (vPFC and dIPFC, respectively), in addition to the ventral medial prefrontal cortex, as critical for regulating amygdala activity [59,66,67,69,71]. In general, studies have shown largely overlapping prefrontal–amygdala activity to suppression and reappraisal strategies, which was characterized by decreased activity of the amygdala and increased activity of the prefrontal cortex – usually including both medial and lateral PFC [3,67]. Furthermore, the frequency of using reappraisal to regulate emotion in everyday life has been shown to be related to decreased amygdala activity, increased prefrontal and parietal activity [72], and greater vmPFC volume [73]. In addition, during an affect labeling task (i.e., putting emotions into words), which can be regarded as a specific form of emotion regulation [66,74], diminished amygdala activity was again associated with greater activity of the mPFC [64,65], vIPFC [66] and also with increased cortical thickness of the vmPFC [74]. These findings provide functional and structural evidence for shared neural mechanisms during different types of emotion regulation strategies.

These neuroimaging findings lead us to an interesting question – does emotion regulation share similar underlying neural mechanisms with more classic forms of cognitive control? Or are there unique brain circuitries recruited by these distinct emotion regulation processes? According to a cognitive control model of emotion regulation [3], the neural representation of emotion regulation can be summarized as interactions between prefrontal (both dIPFC and mPFC) and ACC systems and their influence on subcortical systems, including the amygdala. The two major types of emotion regulation – suppression and reappraisal – have yielded similar results in terms of brain activations, and how the prefrontal and anterior cingulate cortices interact is strikingly similar to other top-down control mechanisms that do not involve emotional processing, such as cognitive control [75–77]. However, another study has shown that emotion regulation through mood-incongruent autobiographical recall recruits the ventral mPFC and vIPFC, but not dIPFC, implying that activations of neural circuitry depend on the type of emotion regulation being used [78]. Taken together, we can tentatively conclude that while emotion regulation does to some extent share similar neural circuitry with cognitive control, it also recruits unique brain regions, such as the vIPFC. Other forms of top-down control processes, such as regulation of appetitive behaviors, attitudes or prejudice, have also shown to use an overlapping amygdala-prefrontal circuitry [79].

Recent findings raise the possibility that a more efficient crosstalk between the amygdala and the prefrontal cortex begets a better ability to regulate one’s emotions. Supporting this idea, the strength of amygdala–mPFC coupling was quantified by computing the functional connectivity between these two areas and comparing this connectivity with how effectively participants regulated their emotions [34]. It was found that the functional coupling between the amygdala–mPFC was strengthened during reappraisal, and that the degree of this functional coupling was positively correlated with the self-reported effectiveness of emotion regulation [34]. A selective increase in the functional coupling of the amygdala with the vmPFC and dIPFC during emotion regulation has also been reported [69], highlighting the importance of efficient communication between the amygdala and the prefrontal cortex in successful top-down control of emotion. Further, functional coupling of the amygdala and vmPFC at rest predicts beneficial outcomes in terms of reported anxiety [11]. Future studies linking the effective success of emotion regulation strategies and the structural and functional connectivity of the amygdala–mPFC circuitry might provide a better understanding of the neural correlates of these emotion regulatory processes.

5. Amygdala–prefrontal circuitry and the interpretation of emotionally ambiguous facial expressions

In humans, patients with selective amygdala lesions have displayed deficits in processing the facial expressions of fear [80], leading to numerous functional neuroimaging studies using presentations of fearful faces to probe amygdala activity [10,81–86]. These studies have shown that the amygdala is particularly responsive to fearful faces compared to other expressions [87], including angry, happy, and neutral [10,81,82,84–86], except for one report [88]. The affinity of the human amygdala for fearful faces compared to these other expressions, provides insights into amygdala function. For example, since the amygdala is more responsive to fearful faces compared to angry faces, which embody a direct threat, it has been suggested that one function of the amygdala is to augment cortical function through the major neuromodulatory centers to assist in the resolution of predictive uncertainty [86,89]. That is, the inherent ambiguity of fearful faces in that they predict the increased probability of threat without providing information about its nature or location – leads to selective activation of the amygdala [86,89].

Given that the amygdala plays a major role in the resolution of predictive uncertainty associated with fearful faces, surprised facial expressions provide a particularly important comparison expression. Indeed, there is evidence that surprise may be the second-most compromised expression in patients with selective amygdala damage, following fear [90]. Fearful and surprised faces have common facial features (e.g., eye-widening), and both expressions indicate the detection of a significant, but unknown, eliciting event [36]. Surprised faces are particularly interesting because, unlike fear, they do not predict the valence of the unknown eliciting event. Indeed, research has shown that surprised faces can be interpreted as either positive or negative in nature [35,36,91]. Previous research has shown that when individuals make valence judgments of surprised faces, ratings reflect individual differences in one’s positivity/negativity bias [35,91] and these differences are mirrored by a distinct pattern of brain activity, which critically involves the vmPFC as well as the amygdala [35]. Specifically, decreased amygdala activity accompanied by increased vmPFC activity was observed in people who interpreted surprised faces as positive, with the reverse brain pattern seen in those who interpreted surprised faces as negative [35]. The role of the vmPFC in resolving the emotional ambiguity (i.e., the valence of a given surprised face) could be understood as a top-down regulatory input to the amygdala, much akin to the neural mechanism of fear extinction or emotion regulation [4]. Indeed, greater vmPFC activity predicts both (a) more positive ratings of surprise and (b) more positive interpretations of an extinguished tone (i.e., tone now predicts no shock) [92]. In a subsequent study [36], positive and negative sentences (e.g., “He just lost $500” or “He just found $500”) were used to provide contextual information for the presented surprised faces, in order to see how brain activity was influenced by
information that provided clear resolution to the source ambiguity problem associated with surprised faces. Again, data from this study showed that greater amygdala response to negative versus positive faces was accompanied by diminished vmPFC activity, and interestingly greater vPFC activity [36]. Thus, similar medial prefrontal-amygdala regions were activated when a context was provided (i.e., valence of the surprised faces were determined by the experimental condition), compared to when the subjects had to judge the valence of the surprised faces themselves – but additional lateral prefrontal regions were recruited in the contextually mediated condition [35].

In summary, data from these experiments collectively suggest that using emotionally ambiguous stimuli such as surprised faces instigates a competition between top-down and bottom-up processes. This engages the amygdala–mPFC circuitry and the balance of activity within this circuit reflects the resolution of the inherent ambiguity of the perceived surprised faces.

6. Amygdala-prefrontal circuitry and anxiety within the normal range

Anxiety is characterized by chronic, nonspecific apprehension and arousal related to the potential occurrence of future threat [93,94]. Neurobiological theories of anxiety have highlighted the central role of the amygdala in the generation and experience of the fear that can give rise to anxiety [48,49], and fear extinction investigations in animals support such theories [48,49]. Similar to the inhibition of previously conditioned fear responses during fear extinction, reduced anxiety is associated with the top-down regulation of amygdala activity by the mPFC [9,65,95]. Findings from anatomical investigations of amygdala connectivity [14,19] and fear extinction studies in animals [56] emphasize the top-down and bottom-up interactions between the amygdala and mPFC regions in anxiety. To put it another way, efficient crosstalk between the amygdala and the mPFC produces a better outcome in terms of controlling anxiety.

Consistent with this framework, a number of functional neuroimaging studies in humans have shown elevated amygdala activity in highly anxious but otherwise healthy individuals [96–100]. For example, increased amygdala activity to unattended fearful faces was associated with higher levels of self-reported anxiety [96], although this effect may be more prominent in women than men [97]. Using backward masking, an experimental paradigm that has been shown to reliably evoke human amygdala activity and mitigate subjective awareness of fearful face stimuli (e.g., [83,85]), it has been reported that increased amygdala activity was linked to elevated anxiety levels [98]. It is worth noting that in these studies, the relationship between increased amygdala activity and anxiety was evident when the subjects were not attending to or were unaware of the stimuli, not when they were attending to or aware of them. This raises the possibility that attention or awareness may be an important factor that interacts with amygdala activation and subsequent reported anxiety. Anxiety was not only associated with elevated amygdala activity to threat-related stimuli (e.g., fearful faces, emotionally negative pictures), but was also associated with increased activity to non threat-related stimuli (neutral faces; [99]), suggesting that amygdala activity may reflect greater anxiety levels even in the absence of clear threat.

Other evidence from the human neuroimaging literature shows that altered mPFC activity is associated with anxiety [5–8,95]. Although changes in mPFC activity has been consistently reported in anxiety research, the spatial location of that activity (i.e., whether it is dorsal or ventral) varies across studies. Depending on the experimental task, different studies have reported divergent results (for review, see [2]) - for example, anxiety reduced activity of the vmPFC in one study [7], and dmPFC in another [5]. More recently, a number of studies have shown that higher levels of anxiety are associated with both decreased vmPFC activity and increased dmPFC activity [6,8], suggesting differential roles for these mPFC subregions in anxiety.

Based on the findings highlighting the importance of both the amygdala and mPFC regions in anxiety, a number of studies have investigated the amygdala–mPFC circuitry in conjunction with anxiety using functional and structural connectivity measures [9–11]. For example, individuals with anxious temperaments had weaker functional coupling between the amygdala and the vmPFC during a task that involved matching fearful and angry faces [9]. Using DTI, it was demonstrated that the structural integrity of an amygdala–vmPFC pathway was compromised in the participants who exhibited high trait anxiety [10]. Furthermore, studies employing resting state functional connectivity methods have shown that the strength of the coupling between the amygdala and mPFC at rest predicted self-reported levels of anxiety ([11,101]), where a positive correlation between the amygdala and vmPFC predicted beneficial outcomes in terms of reported anxiety [11]. In this study the dmPFC showed an opposite relationship to that observed between the amygdala and vmPFC–dmPFC activity at rest that was negatively correlated with amygdala activity predicted lower levels of anxiety. This latter finding is complemented by a recent task-based fMRI study in which amygdala–dmPFC functional connectivity strength was positively correlated with neuroticism (an anxiety-related personality trait characterized by a bias to interpret normal situations as harmful and threatening; [102]) during viewing emotionally negative faces [103]. Taken together, these data suggest that the strength of amygdala–mPFC functional connectivity during rest may represent efficient crosstalk between the two brain regions, which may be responsible for abolishing the generation of anxious states [11]. This idea is consistent with findings from task-based functional connectivity [9]. Findings from these studies all fit well with the idea that efficient crosstalk between the amygdala and the mPFC, perhaps particularly the vmPFC, is critically involved in lowering anxiety levels.

7. Amygdala-prefrontal circuitry and pathological anxiety

Taking individual differences in normal fluctuations in anxiety as our starting point, disrupted bottom-up and top-down emotional and cognitive processes are thought to be a crucial component of symptomology in pathological anxiety. This model suggests an imbalance between the amygdala and the prefrontal cortex, which is typically characterized by hyperactivity of the amygdala and hypoactivity of the prefrontal cortex [104,105].

7.1. Social anxiety disorder

A prevalent subtype of the anxiety disorders [106], social anxiety disorder (SAD) is characterized by intense anxiety during social situations in which the person is exposed to unfamiliar people [107]. To this end, emotional facial expressions provide a particularly useful paradigm for studying SAD, which is thought to involve exaggerated emotional reactivity to social stimuli and the inability to regulate these responses [108,109]. Individuals with SAD reliably showed elevated amygdala reactivity when viewing “harsh” faces (facial expressions displaying anger, contempt, or a combination of both) [110–112], and even neutral faces [113], compared to healthy individuals in fMRI studies. Amygdala reactivity was positively correlated with symptom severity and/or trait anxiety in SAD patients, further demonstrating the neurobiological significance of the amygdala in SAD [111,113,114]. These individuals also had
exaggerated amygdala reactivity to pictures of emotionally negative scenes (i.e., unpleasant and/or aversive) suggesting abnormal neural activity during general emotional, not just social, processing in SAD [114]. Direct examination of neural activity during emotion regulation demonstrated that SAD patients fail to recruit the mPFC [110], implying that the connectivity of the amygdala–mPFC circuitry is disrupted in SAD. A resting state fMRI study showed that SAD patients had markedly reduced functional connectivity between the left amygdala and the medial orbitofrontal cortex [115], corroborating the previous findings assessing a normal range of anxiety [11]. In addition, state anxiety levels in SAD subjects was inversely correlated with the functional connectivity strength between the amygdala and the medial orbitofrontal cortex, further validating the central role of the amygdala–mPFC circuitry in SAD [115]. In addition to these functional abnormalities, SAD patients exhibited compromised structural integrity of the uncinate fasciculus [116], a major white matter fiber tract that is known to connect the amygdala and the orbitofrontal cortex [117]. Each of these studies provides examples of SAD patients’ failure to recruit the proper cognitive regulatory circuits in the brain, and that the functional abnormalities in these circuits may be attributable, in part, to white matter microstructural problems caused by the pathophysiology of SAD.

7.2. Posttraumatic stress disorder

Posttraumatic stress disorder (PTSD) is a stress-induced anxiety disorder characterized by re-experiencing the traumatic event, avoidance of stimuli associated with the trauma, and more generalized symptoms of hyperarousal [107]. PTSD patients show a diminished ability to extinguish this conditioned fear, which may be evidence for prefrontal cortex dysfunction and reduced amygdala inhibition [118,119]. Shin et al. [120] have demonstrated that PTSD is marked by heightened amygdala activation and reduced anterior cingulate and prefrontal cortex activity when viewing fearful faces. Consistent with this finding, PTSD patients exhibited diminished activity in the mPFC to unattended fearful faces [121]. Both studies reported that PTSD symptom severity was associated with decreased mPFC activity, demonstrating the neurobiological importance of this brain region in the pathophysiology of PTSD [120,121]. Likewise, compared to healthy individuals, PTSD patients failed to recruit vmPFC activity when viewing pictures that were threatening, but unrelated to trauma [122]. In patients with PTSD, the default mode network – brain regions that include the mPFC and the posterior cingulate cortex that are believed to be more “active” during rest [123] – has been affected by the pathophysiology of the disorder as well. Specifically, resting state functional connectivity of the posterior cingulate cortex with the perigenual anterior cingulate and the right amygdala is associated with current PTSD symptoms, and that correlation with the right amygdala predicts future PTSD symptoms [124]. Furthermore, supporting these functional studies, there is DTI evidence that the white matter structural integrity of the cingulum bundle is compromised in PTSD patients compared to healthy individuals [125,126]. Therefore, it is clear that not only the functionality of the amygdala and the mPFC are impaired in PTSD, but also their connectivity is disrupted as well.

Future research exploring the similarities and differences between non-anxious, normal anxious, and pathologically anxious individuals is needed. Based on numerous findings highlighting the relationship between the amygdala–mPFC circuitry and anxiety, developing treatments – whether they involve medication or psychotherapy – for anxiety disorders that target these brain regions will prove to be useful.

8. Conclusions

From normal emotion to pathological anxiety, an organism’s reaction to biologically relevant stimuli and the regulation of these responses can be usefully conceived as a constant struggle between bottom-up and top-down brain processes. A wealth of animal and human neuroimaging studies has shown that the amygdala and the prefrontal cortex, particularly the medial regions, are central to these processes. Investigating the connectivity between the amygdala and the prefrontal cortex has provided a deeper understanding of the role of the amygdala–mPFC circuitry in anxiety. Efficient crosstalk between the amygdala and the prefrontal cortex – represented as stronger structural and functional connectivity – predicts beneficial behavioral outcomes in terms of emotion regulation and anxiety.

Acknowledgements

Supported by the National Institute of Mental Health grants to MJK (F31 MH090672) and PJW (R01 MH080716).

References


Shehzad, the Neurosci 2008;18:2735–47.


Margulies MR, Quirk CK, Rajk RM, Quirk RJ, Rajk SL. Recall of fear extinction in humans activates the ventromedial prefrontal cortex and hippocampus in concert. Biol Psychiatry 2007;62:446–54.


Quirk GJ. Memory for extinction of conditioned fear is long-lasting and persists following spontaneous recovery. Learn Mem 2002;9:402–7.


M.J. Kim et al. / Behavioural Brain Research 223 (2011) 403–410 409


