Norepinephrine ignites local hotspots of neuronal excitation: How arousal amplifies selectivity in perception and memory

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Abstract: Emotional arousal enhances perception and memory of high-priority information but impairs processing of other information. Here, we propose that, under arousal, local glutamate levels signal the current strength of a representation and interact with norepinephrine (NE) to enhance high priority representations and out-compete or suppress lower priority representations. In our “glutamate amplifies noradrenergic effects” (GANE) model, high glutamate at the site of prioritized representations increases local NE release from the locus coeruleus (LC) to generate “NE hotspots.” At these NE hotspots, local glutamate and NE release are mutually enhancing and amplify activation of prioritized representations. In contrast, arousal-induced LC activity inhibits less active representations via two mechanisms: 1) Where there are hotspots, lateral inhibition is amplified; 2) Where no hotspots emerge, NE levels are only high enough to activate low-threshold inhibitory adrenoreceptors. Thus, LC activation promotes a few hotspots of excitation in the context of widespread suppression, enhancing high priority representations while suppressing the rest. Hotspots also help synchronize oscillations across neural ensembles transmitting high-priority information. Furthermore, brain structures that detect stimulus priority interact with phasic NE release to preferentially route such information through large-scale functional brain networks. A surge of NE before, during, or after encoding enhances synaptic plasticity at NE hotspots, triggering local protein synthesis processes that enhance selective memory consolidation. Together, these noradrenergic mechanisms promote selective attention and memory under arousal. GANE not only reconciles apparently contradictory findings in the emotion-cognition literature but also extends previous influential theories of LC neuromodulation by proposing specific mechanisms for how LC-NE activity increases neural gain.

Keywords: arousal; attention; emotion; locus coeruleus; long-term consolidation; memory; norepinephrine; perception

1. Introduction

When jolted by a rough skydiving landing, psychologist James Easterbrook observed that his sense of space and time shrank and slowly re-expanded (Easterbrook 1982). This sparked his curiosity about how arousal influences attention. Later he published a review article in which he argued that under arousal, people rely more on central or immediately relevant information and less on peripheral information (Easterbrook 1959). Since his seminal paper, researchers have accumulated many more observations that arousal evoked by emotional events enhances some aspects of perception and memory but impairs others (for reviews, see Mather & Sutherland 2011; Reisberg & Heuer 2004). For example, victims of a crime tend to remember the weapon vividly but forget the perpetrator’s face (Stelley 1992). People also pay attention
to emotional information at the expense of neutral information (Dolcos & McCarthy 2006; Knight et al. 2007). These examples fit with Easterbrook’s formulation that arousal impairs attention to peripheral information. But arousing stimuli can sometimes enhance memory of peripheral neutral information (Kensinger et al. 2007; Knight & Mather 2009). Thus, although it is clear that arousal shapes attention and memory, knowing that something is neutral or spatially peripheral is not enough to predict how it will fare under emotional conditions.

So, then, how does arousal influence the brain’s selection of features to highlight versus suppress? An initial answer to this puzzle was provided by the arousal-biased competition (ABC) model, which posits that arousal does not have fixed rules about which types of stimuli to enhance or suppress. Instead, arousal amplifies the stakes of ongoing selection processes, leading to “winner-take-more” and “loser-take-less” effects in perception and memory (Mather & Sutherland 2011). The ABC model builds on biased competition models proposing that stimuli must compete for limited mental resources (Beck & Kastner 2009; Desimone & Duncan 1995; Duncan 2006). As conceptualized by Desimone and Duncan (1995), both bottom-up and top-down neural mechanisms help resolve competition.

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Bottom-up processes are largely automatic, determined by the perceptual properties of a stimulus, and do not depend on top-down attention or task demands. For example, stimuli that contrast with their surroundings, such as a bright light in a dark room, engage attention automatically even if they are currently goal irrelevant (Itti & Koch 2000; Parkhurst et al. 2002; Reynolds & Desimone 2003). Top-down goals can also bias competition in favor of particular stimuli that otherwise would not stand out. Although not included in the original biased competition models, past history with particular stimuli is also a source of selection bias (Awh et al. 2012; Hutchinson & Turk-Browne 2012). For example, one’s name or a novel stimulus tends to engage attention (Moray 1959; Reicher et al. 1976). In addition, faces, text, and emotionally salient stimuli all grab attention (e.g., Cerf et al. 2009; Knight et al. 2007; MacKay et al. 2004; Niu et al. 2012).

A core aspect of most current theories of visual attention is that these different signals are integrated into maps of the environment that indicate the priority or salience of stimuli across different locations (Itti & Koch 2000; Soltani & Koch 2010; Treisman 1998). Regions in frontoparietal cortex integrating sensory and top-down signals help represent such priority maps (Ptak 2012). Moreover, having both feedforward and feedback connections between sensory regions and cortical priority maps enables distributed representations of prioritized information to modulate their own processing (e.g., lower-level visual features) even further (Klink et al. 2014; Ptak 2012; Serences & Yantis 2007; Soltani & Koch 2010). Thus, priority signals are self-biased to enhance efficient information processing in the brain.

In the ABC model, arousal further biases mental processing to favor high- over low-priority representations, regardless of whether initial priority is determined by bottom-up salience, emotional salience, or top-down goals. Thus, because spatially peripheral information is usually lower priority than central information, arousal usually impairs memory for it (Stebly 1992; Waring & Kensinger 2011). Yet, when peripheral information is perceptually salient or goal relevant, arousal instead enhances memory for it (e.g., Kensinger et al. 2007, Experiment 4). But the ABC model does not tackle how this works in the brain. Previous brain-based models of emotion and cognition also do not account for the dual role of arousal. Most models posit that the amygdala enhances perception and memory consolidation of emotionally salient stimuli, but fail to address how arousal sometimes enhances and sometimes impairs information processing.

In this article we propose the glutamate amplifies noradrenergic effects (GANE) model, in which arousal amplifies the activation difference between high- and low-priority representations via local synaptic self-regulation of the locus coeruleus–norepinephrine (LC–NE) system. According to the GANE model, hearing an alarming sound or seeing something exciting leads to a surge in NE release, which, in turn, enhances activity of neurons transmitting high-priority mental representations and suppresses activity of neurons transmitting lower-priority mental representations. As already outlined, priority is determined by top-down goals, bottom-up factors, and high-level stimulus features (Beck & Kastner 2009; Desimone & Duncan 1995; Fecteau & Munoz 2006).

According to the GANE model, the brain’s primary excitatory neurotransmitter, glutamate, signals priority.
Under arousal, elevated glutamate associated with highly active neural representations stimulates greater NE release, which then further increases glutamate via positive feedback loops. Thus, in these local “NE hotspots,” glutamate signals are amplified. At the same time, wherever NE is released and fails to ignite a local hotspot, inhibitory adrenoreceptors with lower thresholds of activation suppress activity. Higher NE concentration at hotspots also enhances delivery of energy resources to the site of active cognition, synchronizes brain oscillations, and modulates activity in large-scale functional networks. Thus, under arousal, local NE hotspots contrast with widespread NE suppression to amplify priority effects in perception and memory, regardless of how priority was instantiated.

2. Arousal-biased competition in perception and memory

We start by reviewing recent findings supporting Mather and Sutherland’s (2011) ABC model and its novel predictions. Next, we turn to the question of how these arousal effects operate in the brain. A fundamental challenge in understanding how arousal influences cognition is that it sometimes enhances and sometimes impairs information processing. Although most emotion research focuses on how processing of emotional stimuli is enhanced compared with neutral stimuli, emotional arousal can also influence processing of neutral stimuli, and across studies, opposing effects are often seen. How can emotionally salient stimuli sometimes enhance memory for what just happened, but other times impair it? When do arousing stimuli enhance perception and when do they impair perception of subsequent stimuli? Many studies report that emotion increases selectivity (for reviews, see Levine & Edelstein 2009; Mather & Sutherland 2011; Murray et al. 2013), but how do we predict what gets selected?

2.1. Arousal enhances perception of salient stimuli, but impairs perception of inconspicuous stimuli

In previous research on how arousal influences subsequent perception, two types of findings were hard to reconcile. First, arousing stimuli impair perception of subsequent stimuli. For example, people preferentially perceive arousing stimuli (e.g., Anderson 2005; Keil & Ihssen 2004) but fail to perceive or encode neutral stimuli close to arousing stimuli either in time (e.g., embedded in a rapid series of images after an arousing image) (Smith et al. 2006) or in space (Kensinger et al. 2007; Tooley et al. 1987). Second, hearing or seeing an arousing stimulus enhances visual perception of a subsequent Gabor patch (Lee et al. 2014a; Padmala & Pessoa 2008; Phelps et al. 2006).

How can we explain both the enhancing and impairing effects of arousing stimuli on perception of stimuli that appear close in time or space? Initial evidence supports the ABC hypothesis that inducing arousal should have two opposing effects on perception: Arousal should enhance processing of high-priority (more salient) stimuli but impair processing of lower-priority (less salient) stimuli. When asked to report as many letters as they could from a briefly flashed array (Fig. 1), participants reported more of the high-salience letters and fewer of the low-salience letters after hearing an arousing emotionally negative sound than after hearing a neutral sound (Sutherland & Mather 2012). Similar results were obtained when arousal was induced by emotionally positive sounds (Sutherland & Mather, under review). These results indicate that arousal makes salient stimuli stand out more than they would otherwise.

The ABC model also explains the enhanced processing of emotional stimuli, the focus of most previous theoretical accounts (e.g., Kensinger 2004; LaBar & Cabeza 2006; Mather 2007; Murty et al. 2010; Phelps 2004). People tend to prioritize emotional stimuli due to top-down goals (e.g., increasing pleasure and avoiding pain), their emotional saliency (e.g., associations with reward/punishment), and/or bottom-up salience (e.g., a gunshot is loud as well as a threat to safety [Markovic et al. 2014]). Thus, arousing stimuli should dominate competition for representation at their particular spatiotemporal position (Wang et al. 2012).

If the arousing stimulus appears in the exact same location as a neutral stimulus presented less than a second later, it will impair perception of that neutral stimulus, an effect known as emotion-induced blindness (Kennedy & Most 2012; Most et al. 2005). On the other hand, arousing stimuli tend to enhance the dominance of high-priority stimuli that are nearby but not competing for the same

Figure 1. Participants heard an arousing or neutral sound before a letter array was flashed briefly. They then reported as many of the letters as they could. Some of the letters were shown in dark gray (high contrast and, therefore, salient) and some in light gray (low contrast and less salient). Participants reported a greater proportion of the salient letters than the nonsalient letters, but this advantage for salient letters was significantly greater on arousing trials than on neutral trials, and the disadvantage for the nonsalient letters was significantly greater on arousing than on neutral trials (Sutherland & Mather 2012).
spatiotemporal spot. An emotionally salient word that impairs perception of a subsequent target word flashed in the same location 50 or 500 ms later can instead enhance perception of a target word flashed 1,000 ms later (Bocanegra & Zeelenberg 2009), because after the longer interval, the priority of the target word is no longer overshadowed by the emotionally salient word.

2.2. Arousal enhances perceptual learning about salient stimuli but impairs learning about nonsalient stimuli

Interspersing emotional or neutral pictures with a visual search task had opposite effects on perceptual learning of salient and nonsalient targets (Lee et al. 2012). In this study, the targets were always the same, but in one condition they were salient because they differed from the distractors, and in the other condition they were not salient because they were quite similar to distractors. Emotional images enhanced perceptual learning about the salient target lines but impaired learning of nonsalient targets (Fig. 2). Thus, whether arousal enhanced or impaired learning depended on the target’s salience.

2.3. How arousal modulates neural representations depends on salience

A recent study took advantage of the fact that faces and scenes activate distinct representational regions in the brain to test the ABC hypothesis that arousal increases brain activation associated with processing of salient stimuli, whereas it decreases brain activation associated with processing of less salient stimuli (Lee et al. 2014b). On each trial, one yellow-framed face and one scene image appeared briefly side-by-side and then a dot appeared in the former location of one of the images (Fig. 3A). The participants’ task was to indicate the side on which the dot appeared. Participants responded fastest to dots that appeared behind the salient faces on trials preceded by a tone conditioned to predict shock and thereby induce arousal. In a follow-up functional magnetic resonance imaging (fMRI) study, there was an arousal×salience interaction in visual category-specific brain regions, such that arousal enhanced brain activation in the region processing the salient stimulus (i.e., fusiform face area) but suppressed brain activation in the region processing the nonsalient stimulus (i.e., parahippocampal place area) (Fig. 3B) (Lee et al. 2014b).

2.4. Arousal enhances or impairs memory consolidation of representations depending on their priority

So far, we have focused on how arousal enhances processing of subsequent inputs; however, arousal should have similar effects on mental representations currently active at the moment arousal is induced. Previous research has indicated that arousal induced after initial encoding sometimes impairs and sometimes enhances memory of preceding information (Knight & Mather 2009). The critical ABC hypothesis is that experimental manipulation of priority of information should alter the effect of subsequent arousal on memory consolidation.

In the first study testing this hypothesis, participants viewed lists of objects one object at a time, with one perceptual oddball in each list (Fig. 4) (Sakaki et al. 2014a). The oddball was either emotionally salient or neutral. Some participants were asked to recall the name of the oddball picture as soon as the list presentation ended. In this condition, the object shown just before the oddball (e.g., the cabbage in Fig. 4) was low priority. Other participants were asked to recall the name of the object shown just before the oddball (oddball-minus-1 object). Thus, in this condition, the oddball-minus-1 object (e.g., the cabbage) was high priority. After a series of lists, memory for details of all oddball-minus-1 objects was tested. As predicted, positively or negatively emotionally salient oddball pictures enhanced memory for prioritized oddball-minus-1 objects and impaired memory for nonprioritized oddball-minus-1 objects.

Although the brain mechanisms underlying this priority-arousal interaction in memory have yet to be tested, fMRI evidence indicates that arousal enhances activity in regions processing a high-priority stimulus. For example, pairing shock with certain high-priority (i.e., standalone) neutral scenes enhances successful encoding-related activity in the parahippocampal place area (PPA), the brain region specialized to process scene information (Schwarze et al. 2012). Thus, arousal-induced enhancement of brain

Figure 2. Estimated tuning curves for averaged “target” responses as a function of emotion in the high-salience condition (A) and low-salience condition (B). In the high-salience condition, having interspersed emotional pictures enhanced perceptual learning of the exact tilt of the target (55°), whereas in the low-salience condition, emotion impaired learning of the exact tilt of the same target. Figure adapted from Lee et al. (2012).
activity processing prioritized information not only occurs during perception (e.g., Lee et al. 2014b), but also predicts memory for such items.

2.5. Summary

Mather and Sutherland’s (2011) ABC model accounts for both the enhancement and impairment effects of arousal on neutral stimuli across a wide variety of experimental contexts. It makes novel predictions: (1) Arousal before exposure to stimuli should amplify the effects of salience on perception and memory encoding; and (2) Arousal shortly after encoding information should amplify the effects of its goal relevance on memory consolidation. Both effects result from arousal’s differential modulation of representations depending on priority. Other models also highlight the importance of interactions between arousal, attention, and goals (Kaplan et al. 2012; Levine & Edelstein 2009; Montagrin et al. 2013; Talmi 2013). However, so far there has been no account of how arousal amplifies the effects of priority in the brain.

3. Current brain-based models of arousal’s modulatory effects

Before we present our account of how arousal can modulate neural representations differently depending on their priority, we outline how existing brain-based models of arousal and cognition fail to adequately address how arousal has opposite effects depending on representational priority (see Table 1 for an overview).

3.1. Modular vs. “multiple waves” of emotion enhancement in perception

Noticing things like snakes and guns can increase the odds of survival. Consistent with this adaptive importance, emotionally salient stimuli are often detected more rapidly than neutral stimuli (Leclerc & Kensinger 2008; Mather & Knight 2006; Öhman et al. 2001). Explaining the privileged status of emotional stimuli has been the focus of brain models of emotion perception. One common assumption is that the evolutionary value of noticing emotional stimuli led to a specialized emotion module or pathway to evaluate emotional salience (Tamietto & de Gelder 2010). For example, in their multiple attention gain control (MAGiC) model, Pourtois et al. (2013) argue that emotional salience shapes perception via amplification mechanisms independent of other attentional processes. In the MAGiC model, the amygdala and other modulatory brain regions amplify neural responses to emotional relative to neutral stimuli along sensory pathways. The model also posits that these modulations occur parallel to and sometimes in competition with signals from bottom-up

Figure 3. In the functional magnetic resonance imaging study by Lee et al. (2014b), tones conditioned to predict shock (CS+ tones) played before the display of a salient face, and a less salient scene (A) increased activity in the left fusiform face area (FFA) associated with face processing, while decreasing activity in the left parahippocampal place area (PPA) associated with scene processing, compared with tones conditioned not to predict shock (CS– tones) (B). *p < 0.05, **p < 0.005. CS = conditioned stimulus; ISI = interstimulus interval.
(exogenous) and top-down (endogenous) attentional control systems (see also Vuilleumier 2005b).

In contrast, Pessoa and Adolphs (2010) argue against a modular approach to emotion enhancement in perception. In their multiple waves model, affectively and motivationally significant visual stimuli rapidly engage multiple brain sites, including the amygdala, orbitofrontal cortex, anterior insula, and anterior cingulate cortex, that then bias processing to favor these stimuli. From their perspective, the amygdala helps prioritize emotional aspects of information processing by coordinating activity in other regions involved in selective attention. Thus, in the multiple waves model, emotion influences general-purpose perceptual and attention systems rather than harnessing independent brain mechanisms to enhance perception of emotional items.

The latter perspective is more compatible with our findings than are separate-system models; if emotional stimuli were processed via a system separate from that processing neutral stimuli, it is not clear how emotional arousal could have both enhancing and impairing effects on neutral stimuli depending on their priority. However, even this modulatory multiple waves approach to emotion–cognition interactions fails to explain the full picture of how emotional arousal influences cognitive processing, as it focuses only on the enhanced perception of arousing stimuli and ignores how arousal affects perceptual selectivity more generally.

3.2. Canonical amygdala modulation model of emotional memory enhancement

The act of noticing something creates initial trace representations that require additional resources over the next few minutes, hours, and days to consolidate into a longer-lasting memory. Much research indicates that emotional arousal experienced before, during, or after an event can enhance these memory consolidation processes (Hermans et al. 2014). The prevailing view of how emotion affects memory consolidation is that the amygdala enhances processes in the hippocampus and other memory-related brain regions in the medial temporal lobes, such that memory for emotional events is enhanced compared with memory for neutral events (e.g., McGaugh 2004).
Table 1. Brain-based emotion-cognition theories.

<table>
<thead>
<tr>
<th>Models that focus on enhancement of emotionally salient stimuli</th>
<th>Description</th>
<th>Inconsistent/unexplained findings</th>
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<tbody>
<tr>
<td>Multiple attention gain control model (Pourtois et al. 2013)</td>
<td>The amygdala and other modulatory regions amplify emotionally salient signals in the sensory pathway in parallel with bottom-up and top-down systems.</td>
<td>Emotional arousal can enhance perception of not only emotional information, but also non-emotional information.</td>
</tr>
<tr>
<td>Multiple waves model (Pessoa &amp; Adolphs 2010)</td>
<td>The amygdala and other modulatory regions coordinate activity in attention systems to enhance perception.</td>
<td>Emotional arousal does not always enhance perception.</td>
</tr>
<tr>
<td>Amygdala modulation hypothesis (McGaugh 2004)</td>
<td>The amygdala enhances processing in other memory-related regions to enhance memory for emotional events via noradrenergic mechanisms.</td>
<td>Norepinephrine–amygdala interactions enhance memory not only for emotional stimuli, but also for non-emotional stimuli.</td>
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<tr>
<th>Models that address selective effects of emotion</th>
<th>Description</th>
<th>Inconsistent/unexplained findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biased attention via norepinephrine (BANE) model (Markovic et al. 2014)</td>
<td>The anterior affective system detects emotional saliency and recruits the locus coeruleus–norepinephrine system to bias attention and memory in favor of emotionally salient stimuli.</td>
<td>Emotional information sometimes enhances perception and memory for nearby neutral information.</td>
</tr>
<tr>
<td>Dual competition model (Pessoa 2009)</td>
<td>Emotional stimuli compete for resources with other stimuli, leaving fewer resources available for non-emotional stimuli.</td>
<td>Emotional information sometimes enhances perception and memory for nearby neutral information.</td>
</tr>
<tr>
<td>Ruthless competition model (Diamond et al. 2005)</td>
<td>Encoding new emotional information suppresses recently potentiated synapses, resulting in enhanced memory for emotional information at the cost of preceding events.</td>
<td>Emotional arousal enhances memory for what occurred earlier if the preceding event is emotional.</td>
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<tr>
<td>Emotional-tagging hypothesis (Richter-Levin &amp; Akirav 2003)</td>
<td>Memories for emotional events are tagged, which allows for subsequent arousal to selectively enhance memory for preceding emotional events.</td>
<td>Emotional arousal can produce retrograde enhancement even when preceding information is non-emotional.</td>
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Consistent with this idea, activity in the amygdala during encoding predicts later memory for emotional items, but not memory for neutral items, as does greater amygdala functional connectivity with medial temporal brain regions (Dolcos et al. 2004; Kilpatrick & Cahill 2003; Richardsson et al. 2004; Ritchey et al. 2008).

Converging rodent and human research indicates that NE facilitates the amygdala-mediated enhancement of emotional information. For example, NE released in the amygdala during arousal is associated with enhanced memory for the emotionally arousing event (McIntyre et al. 2002). Infusion of noradrenergic agonists into the basolateral amygdala after training also enhances memory for emotionally arousing events (Hatfield & McGaugh 1999; LaLumiere et al. 2003). In humans, administration of the β-adrenergic antagonist propranolol impairs emotional memories, whereas pharmacological agents that increase NE levels, such as a selective NE reuptake inhibitor, tend to enhance them (Chamberlain & Robbins 2013), and enhanced amygdala activity during encoding emotional stimuli is reduced by propranolol (Strange & Dolan 2004). Thus, NE–amygdala interactions enhance memory for emotional events.

Activation of the amygdala by NE can also impair memory for neutral information encountered near something emotional. For example, as already described earlier in the context of the Sakaki et al. (2014a) study, people often have worse memory for neutral-low priority information shown immediately before an emotional compared with a neutral “oddball” stimulus. Patients with amygdalar damage do not exhibit decrements in memory for neutral words preceding emotional oddball words, and in normal individuals, a β-adrenergic antagonist prevents this retrograde memory impairment (Strange et al. 2003).

Although not usually articulated, the amygdala modulation hypothesis presumably explains these impairment effects for neutral stimuli in terms of a trade-off in which the amygdala focuses resources on emotional stimuli, leaving fewer resources available to process and consolidate
the neutral stimuli. However, this trade-off explanation fails to explain how NE–amygdala interactions can also enhance memory for nonarousing information (e.g., Barsegyan et al. 2014; Roozendaal et al. 2008).

3.3. Biased attention via norepinephrine model

In the biased attention via norepinephrine (BANE) model, Markovic et al. (2014) propose that affectively salient stimuli activate the LC–NE system to optimize their own processing. Like the ABC model (Mather & Sutherland 2011), the BANE model builds on biased competition models of attention (Markovic et al. 2014). The BANE model proposes that affect-biased attention "is distinct from both 'classic' executive top-down and bottom-up visual attention and is at least in part circumscribed by a different set of neural mechanisms" (Markovic et al. 2014, p. 230). In the BANE model, emotional salience is detected by an "anterior affective system," including the amygdala and the orbitofrontal cortex, based on the recent history of reward and punishment. In turn, the amygdala’s recruitment of the LC–NE system serves as an additional specialized pathway that further biases attention and memory in favor of the affectively relevant information that triggered NE release. However, like other models of emotion and cognition, the BANE model focuses exclusively on how affectively salient stimuli outcompete less salient stimuli and does not address how arousal induced by these stimuli sometimes enhances and sometimes impairs processing of proximal neutral information.

3.4. Emotional attention competes with executive attention for limited mental resources

Another line of work focuses on how emotional stimuli compete for executive resources (Bishop 2007; Choi et al. 2012; Eysenck et al. 2007), with some researchers positing that a ventral affective system competes with a dorsal executive system (Bush et al. 2000; Dolcos et al. 2011). For example, when task-irrelevant emotional stimuli capture attention, they diminish dorsal executive brain region function and therefore disrupt working memory for neutral faces that were just seen (Dolcos & McCarthy 2006; Dolcos et al. 2008). However, meta-analyses indicate that emotional responses are associated with both the ventral and dorsal prefrontal cortical regions (Phan et al. 2002; Shackman et al. 2011), and so the notion that emotional distractors lead the ventral prefrontal cortical region to inhibit the dorsolateral prefrontal cortical region (Dolcos et al. 2008) is unlikely to be universal across different contexts.

Instead of a ventral/dorsal antagonism model, the dual competition model posits that emotional stimuli compete for resources at both perceptual and executive levels of processing (Pessoa 2009; 2013). For example, when participants heard tones predicting shock, regions within the frontoparietal network were activated (Lim et al. 2009). Recruitment of these regions during intense emotional arousal should make them less available for concurrent neutral task-related processing and lead to behavioral impairments. At the perceptual level of the dual competition model, both cortical and subcortical structures help amplify visual cortex responses to emotional stimuli, again leading to the impaired perception of other concurrent stimuli.

As in the ABC framework, competition is a core feature of these models. These models, however, consider only one type of competition: that between arousing and neutral stimuli/tasks. Critically, our empirical results indicate that arousal also influences competition between two neutral stimuli, such that processing of high-priority stimuli is enhanced, whereas processing of lower-priority stimuli is impaired. It is not clear how, in competition models that focus on competition between arousing and neutral stimuli, arousal would interact differently with low- and high-priority neutral information. For example, such models cannot account for the differential effects of arousing sounds on subsequent perceptually salient versus nonsalient letters (Fig. 1).

3.5. Competition between items for memory consolidation

In a different type of competition account, Diamond et al. (2005) propose that there is “ruthless competition” between novel and existing memory representations, such that encoding a new emotional experience suppresses recently potentiated synapses, creating memory for emotional events at the cost of memory for information learned just before the emotional event (Diamond et al. 2005).

This ruthless competition hypothesis argues that the acquisition of new information via the hippocampus depletes the most recently activated synapses and that this suppression of recently formed memories is greater when the new information induces emotion or stress. Thus, inducing arousal should impair memory for a preceding sequence of items, regardless of whether those preceding items were themselves emotional or not. That is not the case, however. Inducing arousal via emotional or cold-pressor stress immediately after participants study a mixed list of emotional and neutral pictures selectively enhances memory for preceding emotional, but not neutral, pictures (Cahill et al. 2003; Liu et al. 2008).

3.6. An arousing stimulus sometimes impairs and sometimes enhances memory of what just happened

How can inducing arousal enhance memory for preceding emotional items but not neutral items? Investigators proposed that emotional arousal “tags” synapses associated with representations of emotional items, making these synapses the selective target of protein synthesis-dependent long-term potentiation (Bergado et al. 2011; Richter-Levin & Akirav 2003; Segal & Cahill 2009; Tully & Bolsbakov 2010). The emotional tagging hypothesis predicts that emotionally salient stimuli are remembered better than neutral stimuli because emotional tags allow those particular synapses to capture the plasticity-related proteins released with subsequent inductions of arousal.

A problem for the emotional tagging model is that inducing emotional arousal sometimes enhances memory for preceding neutral stimuli (Anderson et al. 2006; Dunsnoor et al. 2015; Knight & Mather 2009; Nielson & Powless 2007; Sakaki et al. 2014a). Neither the emotional tagging hypothesis nor any of the other hypotheses outlined...
earlier can account for this retrograde enhancement of something neutral. In contrast to the emotional tagging hypothesis, behavioral studies demonstrate that whether something arousing will yield retrograde enhancement or impairment depends on the priority of the preceding information (sect. 2.5) (Ponzio & Mather 2014; Sakaki et al. 2014a).

3.7. Summary

Although there are many models describing how emotion enhances perception, attention, and memory in the brain, these theories fail to account for both the enhancing and impairing effects of emotional arousal (see Table 1 for a summary). In the following sections, we make the case for GANE, a model of how NE released under arousal can impact high- and low-priority representations differently despite its diffuse release across the brain.

4. Locus coeruleus, NE, and arousal

Like the GANE model, other theories also argue that the LC–NE system is important for emotion–cognition interactions (Markovic et al. 2014; McGaugh 2000; 2004; McIntyre et al. 2012). However, they have focused mostly on how NE interacts with the amygdala to enhance processing and consolidation of emotional stimuli at the expense of processing neutral stimuli (e.g., Strange & Dolan 2004; Strange et al. 2003). In contrast, we argue that the LC–NE system promotes selectivity for any prioritized stimuli, irrespective of whether they are emotional or nonemotional.

In this section, we review the functional anatomy of the LC–NE system. A small nucleus in the brainstem known as the locus coeruleus (LC) releases NE when people are aroused—which by a reward or punishment, a loud noise, or a disturbing image. LC axons are distributed throughout most of the brain (Gaspar et al. 1989; Javoy-Agid et al. 1989; Levitt et al. 1984; Swanson & Hartman 1975), enabling NE to modify neural processing both locally and more globally in large-scale functional brain networks. How does the LC influence information processing in most cortical and subcortical regions? One might think that a hormone released under conditions of arousal would amp up brain activity. But instead, NE quiets most neuronal activity. In turn, this quiet backdrop makes those select few representations that NE amplifies stand out even more.

4.1. Functional neuroanatomy of the LC–NE system

The LC is the primary source of cortical NE and helps determine arousal levels (Berridge & Waterhouse 2003; Berridge et al. 2012; Samuels & Szabadi 2008a; 2008b). Tonic, or background, levels of LC activity help regulate levels of wakefulness (Carter et al. 2010). Phasic, or transient, bursts of LC activity occur in response to novel, stressful, or salient stimuli (Aston-Jones & Bloom 1981; Foote et al. 1980; Grant et al. 1985; Sara & Bouret 2012; Sara & Segal 1991; Vankov et al. 1995) or to top-down signals associated with decision outcomes or goal relevance (Aston-Jones & Cohen 2005; Aston-Jones et al. 1999). Emotionally salient stimuli also induce LC phasic activity irrespective of whether stimuli are positive (Bouret & Richmond 2015; Grant et al. 1988) or aversive (Chen & Sara 2007; Grant et al. 1988).

With highly divergent branching axons, the LC projects to every major region of cortex, despite its relatively small number of neurons (13,000 per hemisphere in humans) (Foote & Morrison 1987). Subcortical regions that underlie memory, attention, and emotional processing, including the hippocampus, frontoparietal cortex, and amygdala, are also innervated by the LC (Berridge & Waterhouse 2003). LC axon varicosities release NE into extracellular space, allowing it to activate a broad swath of receptors within a diffusion zone (Beaudet & Descarries 1978; Descarries et al. 1977; O’Donnell et al. 2012).

In target brain sites, NE binds to multiple receptor subtypes (i.e., α1, α2, and β receptors) that are located both pre- and postsynaptically on neurons and astrocytes (Berridge & Waterhouse 2003; Marzo et al. 2009; Nomura et al. 2014; Starke 2001). α1-Adrenoreceptors recruit phospholipase activation and typically increase cell excitability by the inhibition of potassium channels (Wang & McCormick 1993). Thus, the relative density and localization of adrenoreceptor subtypes help determine how arousal-induced NE release will affect neural processing in different brain regions.

4.2. NE decreases neuronal noise in sensory regions during arousal

In the 1970s, researchers proposed that LC–NE activity enhances signal-to-noise ratios in target neurons in sensory regions (Foote et al. 1975; Freedman et al. 1977; Segal & Bloom 1976; Waterhouse & Woodward 1980). For example, recording from individual neurons in awake squirrel monkeys revealed that NE application reduced spontaneous activity more than it reduced activity evoked by species-specific vocalizations (Foote et al. 1975). Noradrenergic regulation of signal-to-noise ratios is characterized by two simultaneous effects: (1) most neurons in a population decrease spontaneous firing, and (2) the few neurons that typically respond strongly to the specific current sensory stimuli either show no decrease or an increase in firing, unlike the majority of neurons for which the stimuli typically evoke weak responses (Foote et al. 1975; Freedman et al. 1977; Hasselmo et al. 1997; Kuo & Trussell 2011; Livingstone & Hubel 1981; O’Donnell et al. 2012; Oades 1985; Waterhouse & Woodward 1980).

Intracellular recording data in awake animals support and extend these early observations. Both inhibitory and excitatory neurons are depolarized in aroused cortex when mice run (Polack et al. 2013). Yet, consistent with earlier reports of a quieter cortex under arousal, inhibitory neurons are more depolarized than excitatory neurons (Polack et al. 2013). Moreover, surround inhibition dominates sensory responses during wakefulness compared with anesthesia, increasing the speed and selectivity of responses to stimuli in the center of the receptive field (Haider et al. 2013). NE mediates the increase in widespread
depolarization and the increase in inhibitory activity in visual cortex that together increase the signal-to-noise ratio (Polack et al. 2013). The effect of NE has also been characterized as increasing the gain on the activation function of neural networks (Fig. 5) (Aston-Jones & Cohen 2005).

Arousal is also characterized by cortical desynchronization, both globally when comparing wakefulness with anesthesia (Constantinople & Bruno 2011) or locomotion with being stationary (Polack et al. 2013) and locally among neurons corresponding to attended representations (Fries et al. 2001). Such decreases in cortical slow wave synchrony under arousal are likely mediated by LC activity (Berridge & Foote 1991; Berridge et al. 1993). Synchronous slow wave neural activity may gate sensory inputs, whereas desynchronized activity permits communication of cortical representations of stimuli across the brain (Luczak et al. 2013). Cortical cell depolarization, desynchronization, and increased responsiveness to external input also occur with pupil dilation (Reiner et al. 2014; Vinck et al. 2014), and pupil dilation tracks LC activity (Murphy et al. 2014).

4.3 Summary

Years of research indicate that NE suppresses weak or random neuronal activity, but not strong activity. This is consistent with the increased selectivity seen under arousal (sect. 2). In the next section, we outline a model of how NE has such different outcomes depending on activity level.

5. Glutamate amplifies noradrenergic effects: The core noradrenergic selectivity mechanism under arousal

Now we turn to our GANE model, a novel brain-based account of how arousal amplifies priority effects in perception and memory. We propose that local glutamate–NE interactions increase gain under arousal. Glutamate is the most prevalent excitatory neurotransmitter in the brain (Meldrum 2000). Glutamate receptors such as α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) and N-methyl-D-aspartate (NMDA) receptors mediate rapid excitatory synaptic transmission, neural network connectivity, and long-term memory (Bliss & Collingridge 1993; Lynch 2004; Travénels et al. 2010).

In addition to point-to-point transmission across a synapse, some glutamate escapes the synaptic cleft, resulting in “glutamate spillover” (Okubo et al. 2010). In this section, we outline evidence that glutamate spillover attracts and amplifies local NE release via positive feedback loops. These self-regulating NE hotspots generate even greater excitatory activity in the vicinity of synapses transmitting high-priority representations, in contrast with NE’s suppressive effects in the more widespread non-hotspot regions.

5.1. The NE hotspot: How local NE–glutamate positive feedback loops amplify processing of high-priority information

5.1.1 High glutamate activity stimulates adjacent NE varicosities to release more NE. The first demonstrations of glutamate-evoked effects on NE found that glutamate increased NE release via NMDA and non-NMDA glutamate receptors on LC axons (Fink et al. 1989; Göthert & Fink 1991; Lalleys et al. 1985; Nelson et al. 1980; Pittaluga & Raiteri 1990; 1992; Vezzani et al. 1987; Wang et al. 1992, see also Jones et al. 1987). In these studies, glutamate-evoked NE release occurred for NE varicosities in all cortical structures investigated in vitro: olfactory bulb, hippocampus, and throughout neocortex. In vivo experiments replicated the effect with targeted glutamate in rodent prefrontal cortex (Lehmann et al. 1992). Other neurotransmitters associated with arousal, such as histamine (Burban et al. 2010) and orexin (Tose et al. 2009), enhance glutamate-evoked NE release. Central to our hypothesis, glutamate-evoked NE release occurs in human neocortex (Fink et al. 1992; Luccini et al. 2007; Pittaluga et al. 1999).

How do these glutamate–NE interactions occur? LC axon varicosities rarely make direct synaptic contacts (e.g., only ~5% in rat cortex) (Vizi et al. 2010), but the distribution of these varicosities suggests they should often be found near glutamate terminals at excitatory synapses in neocortex (Benavides-Piccione et al. 2005; Gaspar et al. 1989). Another critical point is that LC neurons produce the NMDA receptor subunits needed for glutamate to modulate the release of NE from LC axon varicosities (Chandler et al. 2014; Grilli et al. 2009; Petralia et al. 1994; Zhu et al. 2003).

New technologies enable the visualization of glutamate spillover in cerebellum, neocortex, and hippocampus (Okubo et al. 2010; Okubo & Iino 2011). Multiple action potentials in a row yield sufficient spillover glutamate to activate nonsynaptic NMDA and group I metabotropic glutamate receptors (mGluRs) (which are co-expressed on NE varicosities and enhance glutamate-evoked NE release in rodent and human cortices [Luccini et al. 2007]), but probably yield insufficient glutamate to recruit lower-affinity AMPA receptors (Okubo et al. 2010). Extracellular concentrations of the spillover rapidly decrease as distance from the synaptic cleft increases (Vizi et al. 2010), and the upper limit of glutamate spillover effects is estimated to be no greater than a few micrometers (Okubo & Iino 2011).

That spillover glutamate is sufficient to activate NMDA, but not AMPA receptors is another key factor. Unlike
AMPA receptors, NMDA receptors require synchronized glutamate stimulation and neuron depolarization to activate (Liischer & Malenka 2012). Thus, local glutamate spillover must co-occur with phasic depolarizing bursts of activity in LC neurons to recruit additional local NE release. Furthermore, a unique feature of NMDA receptors is that they require a co-agonist, which could be either glycine or D-serine (Wolosker 2007). Glutamate stimulates astrocytes to release these co-agonists (Harsing & Matyus 2013; Van Horn et al. 2013), and both glutamate and NE stimulate astrocytes to release glutamate (Parpura & Haydon 2000). These additional glutamate interactions would further enhance NMDA receptor-mediated NE release (Fig. 6) (Paukert et al. 2014). Together, these local glutamate–NE interactions support the emergence and sustainment of hotspots in the vicinity of the most activated synapses when arousal is induced.

Consistent with the existence of glutamate–NE interactions, local NE release in the region of an activated novel representation depends on the coincident timing of the novel event and an arousing event (Rangel & Leon 1995). For example, when footshock was administered to a rat while it explored a novel environment, NE levels rose substantially higher and remained elevated longer than when footshock was administered to the rat in its holding cage (Fig. 7) (McIntyre et al. 2002). The amygdala presumably activated in response to the novelty of the new environment (Weierich et al. 2010), and glutamate associated with that representational network amplified the NE release initiated by the shock.

Hotspot effects have also been observed in the bed nucleus of the stria terminalis immediately after training rats on an inhibitory avoidance task (Liu et al. 2009). When infused separately at low doses, glutamate and NE each had no effect. But when infused together at the same low doses, they produced marked memory enhancements. Infusion of a higher dose of glutamate led to memory enhancements that were blocked by propranolol, indicating that the glutamate effect required β-adrenergic activity, which, as we describe next, is another key feature of our hotspot model.

5.1.2. α- and β-adrenoreceptors exert different effects on neuronal excitability and require different NE concentrations to be activated. To be engaged, β-adrenoreceptors require relatively high NE concentrations, α1-adrenoreceptors more moderate levels, and α2-adrenoreceptors the lowest NE concentrations (Ramos & Arnsten 2007). Thus, under arousal, α2-adrenoreceptor effects should be widespread, whereas β-adrenoreceptors should be activated only at hotspot regions because the local glutamate-evoked NE release there results in higher NE levels. Next, we describe the importance of this distinction for adrenergic autoreceptors.

5.1.3 Adrenergic autoreceptors inhibit or amplify their own NE release. Autoreceptors at NE varicosities serve as neural gain amplifiers by taking opposing action at low and high local levels of NE. The predominant presynaptic noradrenergic autoreceptor in humans is the α2A-adrenoreceptor (Starke 2001), which inhibits NE release when it detects low or moderate levels of NE (Delaney et al. 2007; Gilsbach & Hein 2008; Langer 2008; Starke 2001). In contrast, presynaptic β-adrenoreceptors amplify NE release when activated by high levels of NE (Chang et al. 1986; Misu & Kubo 1986; Murugaiah & O’Donnell 1995a; 1995b; Ueda et al. 1985). In addition, α2A-adrenoreceptors may lose affinity for NE when neurons are depolarized (Rinne et al. 2013), which would remove their inhibitory influence as a region becomes highly active. However, this loss of affinity recovers at saturating levels of NE (Rinne et al. 2013), which should help prevent the runaway excitation that could otherwise emerge because of the NE-glutamate feedback loop. Together with glutamate-evoked NE release (see sect. 5.1.1), the opposing effects of these different autoreceptors at low and high levels of NE provide an elegant way for the LC to modulate signal gain depending on the level of local excitation.

5.1.4 Elevated local NE at hotspots engages β-adrenoreceptors on the glutamate terminals transmitting the prioritized representation. This stimulates an even greater release of glutamate, thereby amplifying the high-priority excitatory signal (Ferrero et al. 2013; Gereau & Conn 1994; Herrera & Sánchez-Prieto 1996; Ji et al. 2008; Kobayashi et al. 2009; Mobley & Greengard 1985). That β-adrenoreceptors require relatively high NE concentrations to
be engaged further biases this form of cortical autoregulation towards the most active synapses. Through these feedback processes, high-priority representations are “self-selected” to produce a stronger glutamate message and excite their connections more effectively under arousal. This stronger glutamate message should also promote selective memory of such stimuli (see sect. 6.1). In contrast, activation of lower threshold α2-adrenoreceptors inhibits glutamate release (Bickler & Hansen 1996; Egli et al. 2005), providing a mechanism for inhibiting lower-priority neural activity under arousal.

5.1.5 Higher NE levels at hotspots help prolong the period of neuronal excitation by temporarily inhibiting processes that normalize neuron activity. Under normal conditions, the slow after-hyperpolarization current habituates a postsynaptic neuron’s responses following prolonged depolarization (Alger & Nicoll 1980). However, even here, NE seems to benefit prioritized inputs by prolonging neuronal excitation via β-adrenoreceptors inhibiting the slow after-hyperpolarization (Madison & Nicoll 1982; Nicoll 1988).

In summary, different receptor subtypes enable NE to ignite hotspots in regions with high glutamate levels while inhibiting activity elsewhere. As we outline later, this diversity in NE receptor subtypes also plays an important role in shaping synaptic plasticity to favor prioritized representations under phasic arousal.

5.2. NE hotspots modulate interneurons and GABAergic transmission to increase lateral inhibition of competing representations

Increases in glutamate and NE at hotspots should also enhance inhibitory activity that mediates competition among neurons. γ-Aminobutyric acid (GABA) is the most widespread inhibitory transmitter from neurons that suppress the responses of other neurons or neuronal circuits (Petroff 2002). Strong glutamate activity in cortical circuits stimulates local GABAergic activity, which increases the inhibitory effects of highly active regions on neighboring, competing neural circuits (Xue et al. 2014). Increases in NE also activate inhibition directly, with intermediate concentrations engaging maximal suppression (Nai et al. 2009).

Subtypes of interneurons respond differently to NE in ways that should further increase neural gain. Although LC–NE activity activates interneurons that mediate lateral inhibition (Salgado et al. 2012a), it can also suppress interneurons with feedforward connections (Brown et al. 2005), such that a strong signal will inhibit competing representations while enhancing activity in other neurons within its processing pathway.

5.3. NE directs metabolic resources to where they are most needed

To optimize processing of salient events, NE also helps coordinate the delivery of the brain’s energy supplies, allowing it to mobilize resources quickly when needed (e.g., Toussay et al. 2013). The brain’s most essential energy supplies, oxygen and glucose, are delivered via the bloodstream. One key way that NE coordinates energy delivery is by increasing the spatial and temporal synchronization of blood delivery to oxygen demand within the brain. For example, in mice, as NE levels increase, overall blood vessel diameter in the brain decreases, but the spatial and temporal selectivity of blood distribution to active task-relevant regions increases (Bekar et al. 2012).

In addition to distributing blood flow, NE also interacts with astrocytes locally to mobilize energy resources throughout the cortex. When a particular area of the brain needs more energy, it can obtain fuel not only from glucose, but also from glycogen in astrocytes (Pellerin & Magistretti 2012). NE speeds up the process of obtaining energy from glycogen (Magistretti et al. 1991; Sorg & Magistretti 1991; Walls et al. 2009). While α1- and α2-adrenoreceptors mediate glutamate uptake and glycogen production in astrocytes, β-adrenoreceptors stimulate the breakdown of glycogen to provide rapid energy support in highly active local regions (O’Donnell et al. 2012), further amplifying NE hotspot activity.

5.4. Summary

At the local neuronal level, NE suppresses most activity, but amplifies the strongest activity as a result of the differential effects of NE on different adrenoreceptor subtypes. The amplification of strong activity occurs via “NE hotspots,” where positive feedback loops between local NE and glutamate release increase the strength of activated representations. To sustain higher levels of activity, hotspots also recruit limited metabolic resources. At the circuit level, the increased glutamate and NE produced at hotspots recruit nearby astrocytes that supply additional energy to active neurons. On a broader scale, NE facilitates the redistribution of blood flow towards hotspots and away from areas of lower activity. Thus, by influencing multiple levels of brain function, NE selectively amplifies self-
regulating processes that bias processing in favor of prioritized information.

6. Roles of the LC–NE system in memory

So far we have focused on how arousal increases the gain on prioritization processes in perception, attention, and initial memory encoding. Now we turn to memory consolidation processes. Experiencing an emotionally intense event influences the vividness and longevity of recent memory traces, enhancing or impairing them based on their priority (e.g., Fig. 4) (Knight & Mather 2009; Sakaki et al. 2014a). Much research has indicated that NE is involved in memory consolidation effects (for a review, see McGaugh [2013]), but there has been little focus on the interplay between NE’s enhancing and impairing effects during memory consolidation.

The durability of memories depends on adjustments in the strength of communication across synapses via processes known as long-term potentiation (LTP) and long-term depression (LTD). Whether neural activity triggers LTP or LTD depends on the relative timing of spikes in pre- and postsynaptic neurons (Nabavi et al. 2014), and whether LTP and LTD are maintained depends on protein synthesis processes (Abraham & Williams 2008). We propose that two main NE mechanisms modulate LTP and LTD, leading to “winner-take-more” and “loser-take-less” outcomes in long-term memory: (1) hotspot modulation of the probability of LTP (higher NE levels engaging LTP) and LTD (relatively lower NE levels promoting LTD), and (2) NE-enhanced protein synthesis supporting long-term maintenance of LTP and LTD.

6.1. NE gates spike-timing-dependent LTD and LTP

Long-term potentiation and long-term depression are often studied in brain slices in a petri dish using high-frequency electric stimulation to induce LTP and repeated slow stimulation to induce LTD. But in the brain’s natural context involving constant barrages of pre-synaptic activity generating postsynaptic spikes, the relative timing of pre- and postsynaptic activity helps determine whether LTP or LTD occurs. Furthermore, to avoid constant up-and-down adjustment of synapses based on random firing patterns, neuromodulators such as NE and dopamine signal when the relationship between pre-synaptic and postsynaptic activity is likely to be meaningful (Pawlak et al. 2010). In vivo studies indicate that spike-timing-dependent LTP or LTD requires these neuromodulators (Huang et al. 2014; Johansen et al. 2014). In particular, by binding to G-coupled receptors, NE modulates kinases and phosphatases that determine whether LTP or LTD induction occurs (Trevisiño et al. 2012b; Tully & Bolskashov 2010).

Different adrenoreceptor subtypes appear to mediate NE’s regulation of spike-timing-dependent LTP and LTD. Spike-timing-dependent LTP is initiated primarily by β-adrenoreceptor activation, whereas α1-adrenoreceptors promote spike-timing-dependent LTD (Salgado et al. 2012b). Critically, Salgado and colleagues reported that the LTD promoting activation of β-adrenoreceptors requires concentrations of NE ~25-fold higher (8.75 µM) than the NE concentration that promotes α1-adrenoreceptor-mediated spike-timing-dependent LTD (0.3 µM) in vitro. This agrees with an in vitro estimate of a 30-fold increase in NE associated with LTP in dentate gyrus (Harley et al. 1996). The increase in NE required to support spike-timing-dependent LTD is substantially higher than the increases in NE levels seen when experimenters stimulate LC and measure NE in cortex or hippocampus using microdialysis (e.g., approximately twice baseline [Florin-Lechner et al. 1996], ~0.5 µM [Palamarchuk et al. 2000]). Thus, there is a discrepancy between the NE levels needed for spike-timing-dependent LTP to occur and the levels measured in laboratory studies. Our GANE model accounts for this difference, as it posits that LC activation interacts with prioritized representations to elicit much higher NE release in a select few local hotspots than elsewhere, such that the average cortical sampling location would not detect the NE levels needed to support LTD.

The NE hotspot model supports a range of simultaneous NE modulatory actions. At high-priority hotspots, NE levels should be sufficiently high to engage β-adrenoreceptors and initiate spike-timing-dependent LTP (Salgado et al. 2012b; Trevisiño et al. 2012b). Conversely, areas with lower glutamate activity, where NE levels are by comparison modestly increased, would undergo LTD as a result of the engagement of relatively higher affinity α1-adrenergic receptors (Huang et al. 2014; Salgado et al. 2012b; Trevisiño et al. 2012a). Variations in NE levels in the alert brain thereby support bidirectional plasticity (Salgado et al. 2012b; Trevisiño et al. 2012b).

6.2. NE increases protein synthesis processes that promote memory consolidation: Critical role of β-adrenoreceptors

Arousal levels in the minutes and hours before or after an event also influence later memory for it. Here we review evidence that these wider time window effects of arousal depend on NE’s enhancement of protein synthesis processes that determine the long-term durability of salient memories. Critically, such regulation of memory processes by NE appears to be mediated by β-adrenoreceptors, which we propose are selectively activated in high-priority representational networks.

The role of NE in gating the synthesis of plasticity-related proteins has been recognized for more than a decade (Cirelli et al. 1996; Cirelli & Tononi 2000). For example, plasticity-related proteins promoted by an LC–NE novelty signal can enhance long-term memory consolidation of another salient, but otherwise poorly consolidated event (i.e., learning that stepping off of a platform leads to a weak shock) that occurs 1 hour later or even 1 hour prior to the novelty experience (Moncada & Viola 2007; Moncada et al. 2011).

Blocking β-adrenoreceptors or protein synthesis prior to novelty exposure prevents novelty facilitation of LTP (Straube et al. 2003). What is particularly striking is that β-adrenoreceptor activation at time 1 primes synapses to induce LTP at time 2 an hour later, even when β-adrenoreceptors are blocked by propranolol during time 2 (Tenorio et al. 2010). However, if protein synthesis processes are blocked during time 2, the time 1 priming event does not lead to enhancement. The plasticity marker, Arc protein, is recruited by β-adrenoreceptor activation in the presence of NMDA receptor activation...
Hotspots are characterized by high levels of glutamate release and β-adrenergic receptor activation; thus, emotional arousal should elevate Arc selectively in NE hotspots.

β-Adrenergic activation after learning or weak LTP induction can also convert short-term LTP to more lasting protein synthesis-dependent late LTP (Gelinas & Nguyen 2005; Gelinas et al. 2008). Likewise, stimulating the basolateral amygdala either before or after tetanization of the hippocampus converts early LTP to late LTP via a β-adrenergic receptor mechanism (Frey et al. 2001). Activation of β-adrenoceptors also shields late LTP from subsequent depotentiation (Gelinas & Nguyen 2005; Katsuki et al. 1997).

Creation of long-lasting memories depends on the protein synthesis cyclic adenosine monophosphate (cAMP)/protein kinase A (PKA)/cAMP response element-binding protein (CREB) pro-signaling cascade (Kandel 2012; O’Dell et al. 2010). Neuronal ensembles in which the cAMP/PKA/CREB cascade has been activated, as happens with the engagement of β-adrenoceptors, have been found to be selectively allocated to the engram representing a memory (Han et al. 2007). Furthermore, increasing excitability via different methods mimics the effects of CREB overexpression, suggesting that neurons are recruited to an engram based on their neural excitability (Frankland & Josselyn 2015; Zhou et al. 2009). Thus, by modulating CREB and other aspects of neural excitability, NE hotspots should help determine which neurons are allocated to an engram and stabilized in long-term memory.

6.3. Summary

Local NE concentration is the key to understanding how NE mediates arousal’s dichotomous effects on memory. Previous research has indicated that different NE levels regulate different forms of spike-timing-dependent plasticity by engaging distinct adrenoreceptors. Whereas NE binding to moderate-affinity α1-adrenergic receptors leads to LTD and memory suppression, NE binding to lower-affinity β-adrenergic receptors leads to LTP and memory enhancement. We propose that local discrepancies in NE levels arise from self-regulating NE-glutamate interactions. Where NE concentrations become high enough to engage low-affinity β-adrenoceptors, a cascade of intracellular events triggers protein synthesis processes that enable long-term memory consolidation of the high-priority trace. In contrast, more modest increases in NE levels at less active regions lead to LTD, ensuring less important events are forgotten. Before or after encoding, the confluence of protein synthesis and β-adrenoceptor activation selectively strengthens memory consolidation when these mechanisms are recruited close in time.

7. Beyond local GANE: Broader noradrenergic circuitry involved in increased selectivity under arousal

Beyond local effects, NE increases biased competition processes by altering how different brain structures interact. With its widely distributed afferents, the LC–NE system influences neural processing in many brain regions when an arousing event occurs. NE release can translate local hotspot effects to more global winnertake-more effects by modulating neuronal oscillations. Furthermore, cortical and subcortical priority signals modulate glutamate release in sensory regions and the hippocampus as mental representations are formed and sustained. As previously reviewed (see sect. 5.1), glutamate is essential for NE release to selectively amplify the processing of significant information. Thus, by stimulating local glutamate release and recruiting LC firing, key brain structures can optimize synaptic conditions for arousal to ignite hotspots.

7.1. Activation of inhibitory networks by NE primes neuronal synchronization among high-priority neural ensembles

So far, we have reviewed evidence that NE hotspots amplify the effects of priority, enhancing salient features while suppressing noisy background activity. In this section, we discuss the possibility that neuronal oscillations communicate activity in local hotspots more globally (Singer 1993).

The first candidate is gamma synchrony (30–80 Hz). Conceptual frameworks of neural oscillations posit that gamma synchrony supports gain modulation in local networks (Fries 2009), such that a target area can oscillate in phase with only one of two competing inputs. As a result, the synaptic input that more successfully synchronizes its activity with the target region is amplified, whereas the less synchronized input is suppressed. Gamma synchrony is likely a key component of selective attention (Baluch & Itti 2011; Fries 2009; Fries et al. 2001).

Gamma oscillations are generated by a feedback loop between excitatory pyramidal cells and fast-spiking parvalbumin-positive inhibitory interneurons (Buzsáki & Wang 2012; Cardin et al. 2009; Carlen et al. 2012; Sohal et al. 2009). Noradrenergic release activates these interneurons (Cox et al. 2008; Huang et al. 2013; Tousay et al. 2013) and increases gamma synchrony in these target regions (Gire & Schoppa 2008; Haggerty et al. 2013; Marzo et al. 2014). Emotional arousal also modulates gamma oscillations in regions that process motivational significance, such as the amygdala, sensory cortex, and prefrontal cortex (Headley & Weinberger 2013). These results suggest that arousal-induced NE release selectively biases gamma oscillations in favor of the most activated representations in local neuronal ensembles.

Consistent with the hotspot model, increases in local gamma power during cognitive processing in humans are associated with increases in glutamate levels (Lally et al. 2014). Increases in local gamma power are also associated with successful memory encoding in humans (Burke et al. 2013). Likewise, in rats, fear conditioning increases gamma synchronization in sensory cortex (Headley & Pare 2013). Increased gamma power predicts retention of tone–shock associations and enhanced representations of the tone associated with shock in the primary auditory cortex (Headley & Weinberger 2011).

Recent research indicates that β-adrenoceptors recruit in-phase oscillations with gamma activity, whereas α1-adrenoceptors recruit out-of-phase oscillations (Haggerty et al. 2013). Given the higher threshold for activating β-adrenergic than α1-adrenergic receptors (see sect. 5.1),
these results suggest that high NE levels at hotspots engage β-adrenoceptors, recruit in-phase oscillations, and increase local network connectivity for prioritized representations. Elsewhere, lower NE levels should only be sufficient to engage α1-adrenoceptors and thereby reduce local gamma power and diminish local synchronization.

In addition to modulating oscillations in local neuronal ensembles, NE also facilitates oscillatory coupling across regions. Current frameworks of neural synchrony posit that long-range/interregional communication between areas is modulated by oscillation in low-frequency bands, such as theta (4–8 Hz), whereas communication within local networks is modulated by high frequencies, including gamma synchrony (Canolty & Knight 2010; Von Stein & Sarnthein 2003). New research further suggests that optimal network function occurs when gamma is embedded in, and phasically facilitated by, slower theta (or even delta [Lakatos et al. 2008]) oscillations (Canolty & Knight 2010; but see Burke et al. 2013). This theta–gamma coupling seems to provide a mechanism for interregional communication and cross-location phase coupling across regions to help translate local NE hotspots into global effects.

Activation of the LC–NE system promotes hippocampal theta (e.g., Berridge & Foote 1991; Walling et al. 2011) and is linked to enhancement of novelty-related hippocampal theta (Koecsis et al. 2007). In humans, the phase coupling of gamma with slower oscillations has been described primarily for neocortex (Canolty et al. 2006), where the role of LC–NE in slower rhythms is less well studied. However, hippocampal theta entrains prefrontal cortical theta (Paz et al. 2008). Recently, selective LC–NE activation was found to increase neocortical theta in anesthetized animals (Vazey & Aston-Jones 2014). The parvalbumin neurons modulated by NE participate in setting not only gamma, but also theta rhythms (Varga et al. 2014; Wulff et al. 2009); thus, parvalbumin interneurons provide a mechanism for LC–NE support of phase-coupled rhythms. Indeed, lesions of NMDA receptors in the parvalbumin neurons result in decreased power of theta oscillations and reduced modulations of gamma oscillation by theta (Korotkova et al. 2010). NE modulation of the hyperpolarization-associated Ih current has also been proposed to support thalamocortical driving of slower neocortical oscillations (Yue & Huguenard 2001). Thus, by modulating gamma and theta, the LC–NE system can amplify the winner-take-more effects of hotspots.

7.2. Key brain regions help evaluate priority and modulate NE hotspots

Here we review how several key brain regions help enhance GANE selectivity mechanisms under arousal. These regions help detect saliency and interact with the LC to fine-tune priority signals via their own hotspot-like effects (e.g., amygdala) and/or other NE mechanisms (e.g., prefrontal cortex and thalamus).

The amygdala plays a central role in enhancing selectivity under arousal. It helps notice and track salient information (Sander et al. 2003) and recruits the LC when activated (e.g., Bouret et al. 2003; Fallon et al. 1978; Jones & Moore 1977; Price & Amaral 1981; Van Bockstaele et al. 1998). The LC, in turn, modulates amygdala activity via NE to further enhance the saliency signal (Sears et al. 2013). Through its strong anatomical projections to sensory cortices (Amaral et al. 2003), the amygdala amplifies cortical processing of behaviorally relevant events (Chau & Galvez 2012; Pessoa & Adolphs 2010). Such modulation of other regions may be mediated by amplification of saliency signals by glutamate–NE interactions within the amygdala (Fig. 7) (see Liu et al. 2009), thereby enhancing the amygdala’s selective modulatory influence on other regions. In addition, as reviewed previously (see sect. 3.2), β-adrenoceptors in the amygdala mediate the selective effects of arousal on memory.

The thalamus helps control the communication of sensory information across the brain (Sherman 2005). Within the thalamus, there are dense NE fibers and high levels of NE in the pulvinar posterior lateral/posterior medial complex, but very few in the lateral geniculate nucleus (Morrison & Foote 1986; Oke et al. 1978). Through its widespread reciprocal connections with cortical and subcortical structures (Shipp 2003), the pulvinar helps filter inputs based on behavioral relevance (Fischer & Whitney 2012), promotes communication across brain regions (Saalmann & Kastner 2009; Saalmann et al. 2012), modulates gamma oscillations (Shumikhina & Molotchnikoff 1999), and controls the gain of sensory processing (Purushothaman et al. 2012). In addition, the pulvinar is sensitive to emotional saliency (Liddell et al. 2005; Padmala et al. 2010; Troiani & Schultz 2013). Thus, anatomically, NE is set up to modulate thalamic signals of priority.

Furthermore, in rats, NE increases signal-to-noise processing within the thalamus. When directly infused with NE, rat ventral posteriomedial thalamus exhibits reduced spontaneous firing, but enhanced firing in response to whisker stimulation (Hirata et al. 2006). When stimulated by phasic or tonic LC activation, ventral posteriomedial thalamus also exhibited increased firing in response to whisker stimulation (Devibiss & Waterhouse 2011). However, an intriguing observation was that in sensory barrel field cortex, phasic stimulation of LC enhanced firing to strong whisker stimulation, but slightly impaired firing to weak whisker stimulation, an outcome consistent with the NE hotspot model. This differential response based on stimulus intensity did not, however, occur within the ventral posteriomedial thalamus, where both strong and weak sensory inputs increased firing (Devibiss & Waterhouse 2011). This initial finding suggests that NE influences in sensory thalamus may occur through mechanisms other than NE hotspots. Thus, further work is needed to examine NE’s modulatory role in the thalamus. In any case, the thalamus plays a key role in amplifying selectivity under arousal by coordinating responses to salient stimuli across the brain. Such local representations of salient stimuli are then subject to NE modulatory influences.

The prefrontal cortex (PFC), including the orbital frontal cortex (OFC) and anterior cingulate cortex (ACC), has reciprocal connections with the LC (Arnsten & Goldman-Rakic 1984; Jodo et al. 1998) and is an important regulator of LC output. PFC regions help appraise sensory information and recruit the LC based on goal relevance (Aston-Jones & Cohen 2005), motivational relevance (Mohanty et al. 2008), reward (for the OFC; Schoenbaum & Roesch 2005), conflict (Botvinick et al. 1999; Sheth et al. 2012), monetary loss (Gehring & Willoughby 2002), and pain (Rainville et al. 1997). The ACC is also a key site for
integrating task-relevant and arousal inputs (Pessoa 2009; Shackman et al. 2011). In humans, LC innervation of the PFC is relatively sparse, especially in anterior regions (Gaspar et al. 1989; Javoy-Agid et al. 1989), but NE modulates working memory processes in PFC (Arnsten 2011; Wang et al. 2007).

These PFC noradrenergic influences on working memory have different mechanisms than the NE hotspot. First, in our model, β-adrenoreceptors support positive feedback loops at NE–glutamate hotspots, but α2-adrenoreceptors suppress those feedback loops (see sect. 5.1). However, the facilitatory versus inhibitory role of these adrenoreceptors reverses in the context of working memory. β-Adrenoreceptors stimulate cAMP, whereas α2-adrenoreceptors inhibit it (Duman & Emna 1986; Nomura et al. 2014; Robinson & Siegelbaum 2003). Inhibition of cAMP via stimulation of postsynaptic α2-adrenoreceptors increases input resistance and enhances recurrent network activity and working memory performance (Wang et al. 2007). Thus, by activating via α2-adrenoreceptors, moderate levels of arousal should enhance working memory processes that maintain goal-relevant information in mind, whereas by activating β-adrenoreceptors, high levels of arousal should impair these processes (Arnsten 2011; Kuhlbandner & Zehetleitner 2011). Such impairments may, in turn, disrupt initiation of top-down prioritization goals after exposure to emotionally salient stimuli (Sutherland et al., in press).

One interesting question is what might occur when top-down priority and bottom-up priority conflict. The insula plays a key role in this aspect and integrates salience signals from internal and external stimuli (Craig 2009; Uddin 2015). The insula is involved in various types of saliency processing, including error detection (Ullsperger et al. 2010), interoception (Craig 2009), oddball detection (Harsay et al. 2012), aversive memory (Miranda & McGaugh 2004), and detection of events that require cognitive resources (Cai et al. 2015). Although not much is known about LC–insula interactions, the LC and other NE brainstem sites project to the insula (at least in rats) Robertson et al. 2013). Neuroimaging studies also suggest that elevated LC–NE activity is associated with encoding-related activity in the insula in response to aversive stimuli (Clerett et al. 2014; Rasch et al. 2009). Consistent with GANE, motivated (higher-priority) versus passive viewing of emotional faces enhances functional connectivity within face processing networks, including the insula and LC (Skelly & Decety 2012; but see Astafiev et al. 2010 for caution when interpreting results from LC (MRI).

7.3. NE amplifies activity in behaviorally relevant functional brain networks

Along with the dorsal ACC, the insula is a key node in a broader “salience network” (Eckert et al. 2009; Hermans et al. 2011) that helps integrate different sources of saliency (Seeley et al. 2007), guide adaptive behavior (Bressler & Menon 2010; Cocchi et al. 2013), and regulate shifts from rest to task-oriented behavior (Sidlauskaitè et al. 2014). On the basis of these findings, recent models of the salience network propose that it mediates competitive interactions between antagonistic attention networks that prioritize internal versus external stimuli (Bressler & Menon 2010; Menon & Uddin 2010). Current data suggest that the LC–NE system modulates salience network activity. For example, β-adrenoreceptor blockade during stress reduces salience network activity (Hermans et al. 2011), and salience network activity is associated with pupil and autonomic responses to errors (Critchley et al. 2005) and overall arousal (Sadaghiani & D’Esposito 2014). In neuroimaging studies, the LC co-activates with the dorsal anterior cingulate during the detection of novel stimuli (Krebs et al. 2013) and during task switching (von der Gablentz et al. 2015), a proposed function of the salience network.

Anatomically, activation of the LC–NE system is well positioned to modulate activity based on priority, as some of the most dense NE innervation is to frontoparietal regions (Gaspar et al. 1989; Javoy-Agid et al. 1989; Morrisson & Foote 1986) that coordinate attention to salient stimuli via priority maps (Ptak 2012). Indeed, phasic LC responses, as indexed by pupil dilation, correlate with activity in a dorsal frontoparietal network during focused attention (Ahnaes et al. 2014). However, more generally, according to the GANE model, activation of the LC–NE system should amplify activity in whichever functional network is currently dominant. Consistent with a role for NE in mediating this process, while subjects rest, pupil dilation increases as activity in the functional network associated with resting state activity increases and activity in a competing motor network activity is suppressed (Yellin et al. 2015). In addition, NE preferentially enhances ventral frontoparietal attention network activity during the detection of salient events that trigger re-orienting (Corbetta et al. 2008; Strange & Dolan 2007). Thus, NE’s influence on gain modulation also manifests at the whole-brain level.

7.4. Summary

Arousal’s dual effects on cognition pervade multilevel brain systems to amplify the priority of important information. By modulating theta and gamma oscillations, NE preferentially synchronizes activity between high-glutamate regions, leading to “winner-take-more” effects in perception and memory. Like some earlier emotion–cognition theories (e.g., Pessoa & Adolphs 2010), the GANE model favors the perspective that the amygdala coordinates information transfer within broader networks that influence salience processing and is not the only route by which NE enhances processing of prioritized stimuli. Brain regions that evaluate saliency modulate LC activity either directly via afferent inputs or indirectly via broader networks. Without contextual signals from these central structures and the periphery, the LC would be blind to salient events that demand attention (Sara & Bouret 2012). In turn, the resulting increase in NE release activates these modulatory structures to further bias neural processing in favor of high-priority stimuli. On a larger scale, NE modulates activity in a salience network that mediates competitive interactions between frontoparietal attention networks supporting higher-level representations of priority. Thus, according to the GANE model, reciprocal interactions between the LC and hierarchical brain networks help strengthen and reinforce priority-biasing signals under phasic arousal (see Fig. 8).
8. Existing models of LC modulation of cognition

In this section, we discuss how the GANE model relates to existing theories of LC neuromodulation of cognition that we have not already discussed.

8.1. Adaptive gain theory

The adaptive gain theory (Aston-Jones & Cohen 2005) posits that two different modes of LC activity (phasic vs. tonic) adaptively adjust the gain of cortical information processing to optimize behavioral performance. Phasic LC activity serves as a temporal attentional filter to selectively process task-relevant stimuli and filter out task-irrelevant stimuli, whereas tonic LC activity regulates overall arousal level in the brain. Phasic LC responses to target detection are constrained by background LC activity and occur most frequently during moderate levels of tonic activity (Usher et al. 1999). Adaptive gain theory provides predictions similar to those of the GANE model in terms of the role of the phasic LC mode: phasic LC activity should increase the gain of task-relevant inputs over noisy or task-irrelevant activity. Our GANE model provides a neuromechanism for these effects by proposing that low to moderate NE levels create ideal conditions to ignite and sustain local NE hotspots via greater phasic LC responses. In support of this notion, a recent fMRI study used baseline pupil dilation before trials of a reward-learning task as a measure of tonic LC–NE activity (Eldar et al. 2013). Both low baseline pupil diameter before the trial and high pupil dilation response during the trial were associated with stronger brain activation in response to task-relevant, but not task-irrelevant stimuli.

8.2. Network reset theory

The LC–NE system activates in response to various salient stimuli, including novel, uncertain, or emotionally salient stimuli (Sara 2009; Yu & Dayan 2005). The network reset theory proposes that when these stimuli are detected, the LC issues a phasic "reset" signal that reorganizes neural networks to facilitate behavioral and cognitive shifts accordingly (Bouret & Sara 2005; Sara & Bouret 2012). This theory explains why emotionally salient stimuli and the sudden onset of goal-relevant or perceptually salient stimuli are preferentially perceived and remembered: these events activate the LC, which then reconfigures functional brain networks to process new sources of priority while impairing ongoing processing of other stimuli. This model, however, does not offer a clear explanation of why phasic arousal induced when encountering emotional stimuli can enhance processing of preceding stimuli when they have high priority.

To explain both the facilitative and impairing effects of emotional arousal on preceding stimuli, the GANE model posits that the incidental release of NE by something emotional can instead maintain–or even enhance–ongoing functional network connectivity when those networks are...
highly activated. Stimulating the LC can inhibit feedforward inhibition by interneurons, thereby increasing the throughput of coincident sensory (glutamatergic) inputs (Brown et al. 2005). Although this “loosening” of neurotransmission enables network flexibility and the building of new representations, the GANE model’s prediction that strong glutamatergic signals transmitting a prioritized representation will benefit from sudden LC activation explains how the “reset” signal triggered by phasic LC activity can still enhance processing of preceding high-priority stimuli.

8.3. Summary

The GANE model both complements and extends previous models of how cognition is influenced by the LC–NE system. According to adaptive gain theory, high phasic LC activity promotes exploitation of the current focus of attention over exploration of other options. In contrast, the network reset theory proposes that phasic LC activity promotes a global reset of attention. The GANE model reconciles these two theories by highlighting the role of priority. According to the GANE model, if the current focus of attention has sufficient priority to yield high glutamate release in synapses transmitting those stimuli, then a phasic increase in LC activity should enhance processing of those representations. Otherwise, increases in LC activity should shift attention and neural resource allocation towards new sources of priority.

The GANE model extends current models of LC function by positing that under arousal, local glutamate–NE interactions will amplify activity of high-priority representations regardless of how those representations initially became highly active. Thus, although the GANE model provides neural mechanisms that account for arousal increasing biased competition outcomes, it can also accommodate other models or modes of information prioritization (Friston 2010; Keitel et al. 2013; Reynolds & Heeger 2009; Wieser et al. 2011).

9. Potential boundary conditions and questions for future research

In this article, we have argued that arousal leads to winner-take-more and loser-take-less effects in perception and memory via local and global noradrenergic mechanisms in the brain. Yet, although the GANE model explains many findings observed in the emotion–cognition literature, there are a number of important questions for future research.

Arousal may not increase selectivity as effectively among older adults because of age-related changes in the LC–NE system, including loss of LC neurons (Manaye et al. 1995; Mather & Harley 2016; Sladek & Sladek 1978; Vijayashankar & Brody 1979). Recent autopsy evidence indicates that lower LC neuron density is related to the rate of cognitive decline prior to death, even after controlling for decline in other aminergic nuclei (e.g., dorsal raphe, ventral tegmental area) (Wilson et al. 2013). β- and α2-adrenoreceptors may also be affected in aging (e.g., Bigham & Lidow 1995; Kalaria et al. 1989). Decreases in α2-adrenoreceptor activity may contribute to age-related cognitive declines because agonists that engage α2A-adrenoreceptors can improve age-related deficits in working memory (Arnsten & Cai 1993; Arnsten & Goldman-Rakic 1985; Ramos et al. 2006), potentially via α2A-induced improvements in the ability to maintain focused attention (Decamp et al. 2011). Aging also affects how effectively glutamate triggers additional NE release (Gonzales et al. 1991; Pittaluga et al. 1993), which would disrupt the emergence and/or efficacy of NE hotspots in older adults.

Another question involves sleep, which plays a crucial role in selectively consolidating salient memory traces (Dieckmann & Born 2010), including emotional stimuli (Hu et al. 2006; Payne et al. 2008; 2012) and top-down prioritized information (Rauchs et al. 2011; Saletin et al. 2011). Emerging research suggests that the LC–NE system may enhance memory consolidation during slow wave sleep (non–rapid eye movement [NREM]), a period when high-priority neural ensembles reactivation (for a review, see Dang-Vu et al. 2008; Eschenko et al. 2012; Sara 2010). For example, a learning-dependent increase in LC activity occurs during slow wave sleep (Eschenko & Sara 2008), and depleting NE prior to encoding reduces slow wave sleep that night (Cirelli et al. 2005). Pharmacologically enhancing LC–NE system activity during slow wave sleep improves recognition of odors learned within the previous 3 hours, whereas blocking LC–NE activity impairs odor recognition (Gais et al. 2011). Blocking NE during sleep also leads to greater memory impairment for emotional than for neutral stimuli (Groch et al. 2011). The timing of transient LC activity coincides with the slow wave grouping of hippocampal sharp wave ripple complexes and sleep spindles that promote NMDA-mediated cellular plasticity (Dieckmann & Born 2010; Rosanova & Ulrich 2005). NE may interact with these processes, given evidence that pharmacological activation of β-adrenoreceptors facilitates the emergence of sharp waves and the induction of LTP (Ul Haq et al. 2012). Together these findings raise the intriguing possibility that the precise timing of NE release interacts with the reactivation of high-priority memory networks to facilitate GANE effects during slow wave sleep.

In this article, we focused on perception, encoding, and consolidation processes, but another important question for future research is how NE modulates memory retrieval (e.g., Sterpenich et al. 2006). For example, when encountering a new experience, our memory system can either store this novel information as a distinct memory (i.e., requiring pattern separation) or use it to reactivate existing memories (i.e., requiring pattern completion) (Bakker et al. 2008). Previous research indicated that arousal facilitates pattern separation (Segal et al. 2012) and that NE facilitates retrieval or pattern completion (Devanges & Sara 1991). But it has been unclear how NE/arousal modulates competition between these two hippocampal processing modes. Glutamate amplification of noradrenergic effects might also affect the stability of a salient memory after it is retrieved, or reconsolidated, because this process involves β-adrenoreceptor and NMDA receptor activation (Lee et al. 2006; Przybyslawski et al. 1999).

Another open question concerns the timing of these effects. Behavioral data indicate that presenting an emotionally salient item influences memory of items appearing in the past few seconds (e.g., Sakaki et al. 2014a) and memory of items appearing in the next few seconds, as well (e.g., Sutherland & Mather 2012). It is plausible that the phasic release of NE would have effects on this time scale, but research examining NE–glutamate interactions is needed to address this question.
On the tonic side of the equation, events that induce stress activate both the LC–NE system and the hypothalamic pituitary adrenal (HPA) axis (Pacak & Palkovits 2001; Sved et al. 2002), and these two systems interact in many ways, especially via the actions of corticotropin releasing factor (CRF). Released by the hypothalamus under stress, CRF helps to initiate the HPA axis response while also targeting the LC (Carrasco & Van de Kar 2003; Valentino & Van Bockstaele 2001; Van Bockstaele et al. 2001). CRF influences both tonic LC activity and sensation-induced phasic discharge, either enhancing or impairing sensation-evoked phasic responses depending on waking state and CRF levels administered (Bangasser & Valentino 2012; Devilbiss et al. 2012; Zitnik et al. 2014). One possibility is that by modulating tonic levels of LC activity, stress also enhances or constrains the impact of phasic arousal responses (see sect. 8.1).

Human genetic studies suggest that different NE polymorphisms moderate the strength of arousals’ influence on memory and perceptual processing. To date, much of this research has focused on the ADRA2B deletion variant in which there is reduced NE inhibitory signaling. In human ADRA2B deletion carriers, there is greater activity in the amygdala and insula during the viewing or encoding of emotional versus neutral images (Cousijn et al. 2010; Rasch et al. 2009). Such patterns of NE-related activity are believed to underlie the larger advantage of emotionally salient over neutral images (Cousijn et al. 2010; Rasch et al. 2009). Such interactions may allow for more nuanced effects and phasic arousal responses depending on emotional input, or some other factor.

It is, however, unclear how these genetic effects relate to the NE hotspot mechanisms outlined in the GANE model. Whereas α2A-adrenoreceptors are found throughout much of the brain and have been clearly identified as autoreceptors regulating NE release, the α2B-adrenoreceptor associated with this genetic polymorphism have a different profile (Brede et al. 2004). They are most dense in striatum, globus pallidus, and thalamus (De Vos et al. 1992; Saunders & Limbird 1999) and are essential for regulating the fetal blood supply (Brede et al. 2004). Thus, although it is possible that these genetic effects alter the feedback cycle in NE hotspots, the genetic differences could also be mediated by different developmental pathways, thalamic modulation of emotional input, or some other factor.

Related to this point about the differential brain localization of α2B-adrenoreceptors is the more general question of how regional variation in receptor density (e.g., Zilles & Amunts 2009) modulates hotspot effects. Modeling and direct comparisons of NE–glutamate interactions across regions could help address this question. In addition, although we have focused on how the LC–NE system influences cognition, other neuromodulators such as serotonin, dopamine, and acetylcholine share many mechanisms of action with NE (Hurley et al. 2004) and interact with NE to regulate attention, memory, and arousal (Arnsten 2011; Briand et al. 2007; Sara 2009). Such interactions are likely to modulate the NE–glutamate interactions highlighted here (some examples already described in sect. 5.1 are interactions with orexin, histamine, glycine, and serine). These interactions may allow for more nuanced effects and some redundancy within the arousal system. However, given NE’s core role in arousal and broad innervation of much of the brain, including source nuclei of other neuromodulators (e.g., ventral tegmental area and basal forebrain) (Jones 2004; Sara 2009), we expect that it plays the lead role in modulating cognitive selectivity as arousal levels fluctuate.

10. Conclusion

Selection is at the core of what allows our cognitive systems to function effectively, enabling us to process the constant influx of information and retrieve the experiences most relevant for adaptive behavior and maintenance of well-being. The ability to focus on salient information is especially important during situations that induce arousal, such as during exposure to threatening or exciting sounds or objects and the pressure to perform a challenging task. For more than 50 years, there has been robust behavioral evidence that arousal often simultaneously enhances and impairs processing of different types of neutral information (Easterbrook 1959). Yet brain-based accounts of how arousal influences cognition failed to address how such dual effects could arise.

Our GANE model fills this critical gap. In this framework, we propose that increases in NE levels under arousal enhance the selectivity of information processing. GANE builds on the previous ABC model (Mather & Sutherland 2011) to provide neural mechanisms of how NE leads to winner-take-more and loser-take-less effects in perception, attention, and memory. Unlike the ABC model, however, the GANE model does not require competition to be a fundamental mechanism. Instead, the GANE model selectively amplifies the activity of whatever priority mechanisms are operating.

Under phasic arousal, local glutamate signals corresponding to a highly activated percept interact with NE to create a hotspot of even higher levels of activity, whereas lower-priority representations are either neglected or further suppressed. These self-regulating hotspots are further aided by NE’s recruitment of brain structures and large-scale functional networks that determine which stimuli deserve attention. NE directs blood flow and energetic resources to brain regions transmitting prioritized information. It supports selective memory consolidation via initiation of LTP and LTD. Through all of these processes, NE increases the gain of prioritized information in the brain, such that things that matter stand out even more and are remembered even better, while the mundane and irrelevant recede even farther into the background and are ignored or forgotten.

Open Peer Commentary

Glutamate and norepinephrine interaction: Relevance to higher cognitive operations and psychopathology

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Abstract: Mather and colleagues present an impressive interdisciplinary model of arousal-induced norepinephrine release and its role in selectively enhancing/inhibiting perception, attention, and memory consolidation. This model will require empirical investigation to test its validity and generalizability beyond classic norepinephrine circuits because it simplifies extremely complex and heterogeneous actions including norepinephrine mechanisms related to higher cognitive circuits and psychopathology.

In their target article, Mather and colleagues propose a molecular model, glutamate amplifies noradrenergic effects (GANE), through which arousal enhances or inhibits perception, attention, and memory consolidation. In this model, arousal precipitates phasic release of norepinephrine (NE) throughout the brain, but “hotspots” of NE release are generated near activated glutamate circuits, sufficient to engage low-affinity β-adrenoceptors, which further increase glutamate release and enhance postsynaptic plasticity by increasing cAMP signaling. The model provides an impressive integration across several fields but will require empirical investigation to test its validity. Furthermore, although the model is presented as universally applicable throughout the brain, NE actually has very heterogeneous actions in different brain circuits. In particular, although the GANE model addresses the effects of normal arousal mechanisms in sensory cortex and hippocampus, it is important to discuss how this model may relate to NE actions in higher cognitive circuits and to conditions of psychopathology.

The noradrenergic system plays an essential role in the pathophysiology and treatment of psychiatric disorders. For example, noradrenergic dysregulation is associated with post-traumatic stress disorder (PTSD), and α1-agonists can reduce these symptoms (Arnsten et al. 2015b; Southwick et al. 1999). Many antidepressants target the noradrenergic system (Klimek et al. 1991), and α1-agonists enhance cognition in patients with attention deficit hyperactivity disorder (ADHD) (Arnsten & Wang 2016). Similarly, accumulating evidence implicates glutamate in the etiology and treatment of mental disorders (Chambers et al. 1999; Krystal et al. 2013). Whether the GANE model applies to traumatic stress conditions is not clear; the research Mather et al. cite utilized subtle arousing conditions—e.g., an emotional word. It is, however, likely to explain several aspects of PTSD; for example, enhancement of the consolidation of traumatic events that may contribute to flashbacks and intrusive memories. However, additional, higher brain changes during trauma may not be captured by this model, as NE actions in the brain are more heterogeneous than described.

Most important for human cognition, the newly evolved circuits in layer III of the dorsolateral prefrontal cortex (dlPFC) that underlie higher cognitive operations are modulated in a unique manner that is often opposite to that of classic synapses in sensory cortex, amygdala, and hippocampus (Arnsten et al. 2012). Indeed, these newly evolved “delay cell” circuits in the dlPFC are even regulated differently than sensory/response-related neurons within the dlPFC. For example, delay cell persistent firing is mediated by NMDAR with NR2B subunits that are exclusively in the postsynaptic density, not extrasynaptic as they are in classic synapses (Wang et al. 2013). Furthermore, delay cells are only subtly influenced by AMPA receptors and show reduced, rather than increased, neuronal firing following systemic ketamine (Wang et al. 2013). In contrast, response feedback cells in the dlPFC (likely layer V) have a more classic profile, with large AMPA receptor influences and show increased, rather than decreased, neuronal firing following systemic ketamine (Wang et al. 2013). These marked differences extend to intracellular cAMP signaling events as well. In classic synapses, activation of cAMP signaling, for example, arising from β-adrenergic receptor stimulation, increases glutamate release from axon terminals and strengthens long-term potentiation (LTP) postsynaptically. However, in layer III dlPFC circuits, increased cAMP signaling weakens connections by opening cAMP-PKA-regulated potassium channels in dendritic spines (Arnsten 2015; Arnsten et al. 2012). Instead, it is inhibition of cAMP signaling via postsynaptic α2A-adrenoceptors that strengthens network connectivity by closing potassium channels near the synapse (Wang et al. 2007). There is currently no evidence of NE “hotspots” in these circuits; for example, blockade of β-receptors within the primate dlPFC has no effect on working memory performance (Li & Mei 1994), even though there are likely high levels of glutamate release in dlPFC arising from the persistent firing of these neuronal networks. Thus, the model in Figure 6 of the target article is misleading because it does not differentiate NE actions in classic synapses from those in more newly evolved dlPFC circuits.

Mather et al. also provide an oversimplified discussion of NE actions at α1-adrenoceptors. Although they focus on α1 mechanisms that weaken plasticity, α1 promotes synaptic actions in many synapses—for example, in somatosensory cortex (Mouradian et al. 1991; Waterhouse et al. 1981; 2000). There are also key circuits where α1-receptor activation potentiates β-receptor actions: For example, in amygdala, α1-receptors facilitate β-adrenergic enhancement of memory consolidation (Ferry et al. 1999a; 1999b). These effects are opposite those described by Mather and colleagues. Their model also does not capture the important finding that high levels of NE release in PFC during stress decrease persistent firing and working memory abilities through stimulation of α1-receptors (Birnbaum et al. 2004). All of these actions likely have a key effect in switching control of behavior from thoughtful, flexible, top-down control by PFC under conditions of safety (moderate levels of arousal) to reflexive, unconscious habits mediated by sensorimotor cortex and subcortical structures during uncontrollable stress (very high levels of arousal).

These mechanisms have particular relevance to the symptoms of PTSD, for which there is extensive evidence of elevated noradrenergic activity (Southwick et al. 1999). For example, the α1-antagonist yohimbine worsens symptoms and induces hypofrontality in subjects with PTSD at doses that have little effect in control subjects (Brenner et al. 1997; Southwick et al. 1993). These drug actions may arise from a combination of neural events, for example, loss of dlPFC top-down control from blockade of postsynaptic α2A-receptors and increased NE stimulation of α1-receptors in dlPFC, as well as increased NE release in “hotspots” in the amygdala, hippocampus, and sensory cortex that may exacerbate anxiety and flashbacks (Arnsten et al. 2015b). Thus, the GANE model may apply to NE actions in classic brain circuits, but not to those in higher cortical circuits, which are strengthened by α2A— rather than β-adrenoceptor mechanisms.

Why we forget our dreams: Acetylcholine and norepinephrine in wakefulness and REM sleep

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Abstract: The ascending fibers releasing norepinephrine and acetylcholine are highly active during wakefulness. In contrast, during rapid-eye-movement sleep, the neocortical tone is sustained mainly by acetylcholine. By comparing the different physiological features of the norepinephrine and acetylcholine systems in the light of the GANE (glutamate amplifies noradrenergic effects) model, we suggest how to interpret some functional differences between waking and rapid-eye-movement sleep.
Regulation of neocortical circuits by ascending regulatory systems involves all of the classic neurotransmitters. Most of the nuclei located in the brainstem, hypothalamus, and basal forebrain not only are reciprocally connected, but also send direct projections to the neocortex (Jones 2011; Saper et al. 2010; Steriade & McCarley 2005). The same applies to release by the hypothalamic nuclei of neuropeptides such as orexin/hypocretin in wakefulness and melanin-concentrating hormone (MCH) in rapid-eye-movement (REM) sleep (Araçri et al. 2015; Jones & Hassani 2008; Monti et al. 2013; and references therein). As a first approximation, these bewildering intricacies can be simplified by focusing on the balance in activity between noradrenergic and cholinergic nuclei, which are crucial regulators of arousal and cognition (e.g., Constantinople & Bruno 2011; Schmidt et al. 2013). Both project varicose fibers that widely innervate the neocortex, and their global effects are excitatory. During wakefulness, high levels of norepinephrine (NE) and acetylcholine (ACh) cooperate in regulating arousal and cognitive processes. However, although cholinergic transmission is certainly implicated in synaptic plasticity (e.g., Constantinople & Bruno 2011), the physiological action of NE is thought to be more persistent and more closely related to memory retention (e.g., Constantinople & Bruno 2011; McGaugh et al. 2010; Takahashi et al. 2010) (Fig. 1). The fact that neocortex activation in REM sleep is sustained mainly by ACh is a further indication that the cholinergic tone is more directly related to consciousness and executive functions. In fact, the role of REM sleep in memory consolidation remains controversial (Ackermann & Rasch 2014; Rasch & Born 2013).

Does the GANE model help suggest possible explanations of the different functional consequences of activating these regulatory systems during brain states? A first central assumption is that, under strong neuronal activation, spillover glutamate stimulates nearby NE varicosities in an N-methyl-D-aspartate (NMDA) receptor-mediated manner. By activating low-affinity β-adrenoreceptors, high NE release would stimulate neuronal excitability, as well as glutamatergic terminals, thus constituting activity “hotspots” that effectively amplify inputs with high priority under phasic arousal. Are such hotspots possible in the cholinergic system? Not much is known about the glutamatergic regulation of ACh release, but evidence does exist of ionotropic glutamate receptors regulating cholinergic terminals in the neocortex (Ghersi et al. 2003; Farikh et al. 2008). Hence, it is conceivable that spillover glutamate also stimulates cholinergic fibers. Because it is well known that AChs increases glutamate release (Marchi & Grilli 2010), a positive feedback loop could generate local ACh hotspots, analogous to those hypothesized by Mather and colleagues.

A second tenet of the GANE model is that the low-threshold α2-adrenoreceptors, by responding to low NE concentrations, would inhibit glutamate release in pathways implicated in low priority signaling, under aroused conditions. In this respect, the cholinergic system presents several differences compared with the noradrenergic. In particular: (1) cholinergic fibers form both well-differentiated point-to-point synapses and axon varicosities that sustain diffuse ACh release (Dani & Bertrand 2007); and (2) ACh activates both metabotropic (muscarinic, mAChRs) and ionotropic (nicotinic, nAChR) receptors. In prefrontal regions, M1 mAChRs are widespread and produce excitatory effects related to working memory through different cellular mechanisms (e.g., McCormick & Prince 1986; Gulledge et al. 2009; Proulx et al. 2014). Their EC50 for ACh is in the low μM range. On the other hand, nAChRs can be divided into two functional classes (Dani & Bertrand 2007). Heteromeric nAChRs have high affinity for ACh (with EC50 in the μM range), relatively low permeability to Ca2+ (Pca), and slow desensitization in the presence of agonist. Homomeric nAChRs have high Pca (in the order of the one displayed by NMDA receptors), but low affinity for ACh (EC50 ≈ 200 μM), and quick desensitization kinetics. A striking difference with NE transmission is immediately apparent. The long-term effects on synaptic consolidation are thought to depend on Ca2+ signals. However, within the putative ACh hotspots, the efficacy of high-Pca homomeric receptors would be blunted by quick desensitization. High ACh concentrations would also tend to desensitize heteromeric nAChRs. This would prevent sustained Ca2+ entry through nAChRs as well as by nAChR-dependent activation of glutamate release, and thus of NMDA receptors. Therefore, it seems unlikely that ACh hotspots can produce long-term cellular effects considerably different from those produced by lower ACh concentrations.

In summary, by following up the GANE model reasoning, one is led to conclude that low and high concentrations of NE and ACh produce distinct functional effects on neocortical networks. Low to moderate ACh release sustains global neocortex arousal in both wakefulness and REM sleep. However, in the absence of NE activity (as in REM sleep), cholinergic activity is unable to yield long-term synaptic changes, such as those implicated in memory retention, which would partly explain the well-known difficulty of recalling oneiric activity. Instead, high levels of ACh seem more able to shape the rapid synaptic responses implicated in executive functions, as the quick kinetics of the low-affinity nicotinic ACh receptors would suggest. We believe that deeper functional studies of the interplay between the ascending regulatory systems, led by heuristic models such as GANE, will greatly lead to progress in understanding the physiological basis of cognition.

For better or worse, or for a change?

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Abstract: The noradrenergic system is intimately related to the autonomic system and is thought to play a key role at the interface between arousal and cognition. The GANE (glutamate amplifies noradrenergic effects) theory proposes a complete account of that role, with an emphasis on the qualitative effect of noradrenaline on stimulus processing. This is in marked contrast to network reset theory, which emphasizes the qualitative effect of noradrenaline of updating the representation of the environment.

Among all neuromodulatory systems, the noradrenergic system is probably the one most closely related to vigilance and autonomic arousal (Aston-Jones et al. 1991; Berridge & Waterhouse 2003; Carter et al. 2010; Foote et al. 1980; Jacobs 1986; Sara & Boulou 2012). The activity of locus coeruleus (LC) neurons is so closely related to arousal that autonomic measures such as pupil diameter are used as a proxy for LC activity in human studies (Einhauser et al. 2008; Jepma & Nieuwenhuis 2011; Nassar et al. 2012; Preuschlo et al. 2011; Sterpenich et al. 2006; Varazzani et al. 2015). Much less clear, however, are the nature of the influence of LC activation–noradrenaline (NA) release on its targets in the brain and its implication for cognition. The GANE (glutamate amplifies noradrenergic effects) theory developed by Mather et al. is addressing this issue directly. This theory covers many aspects of cognitive function, ranging from attention and decision making to memory and emotions, and proposes an original cellular mechanism.

In line with earlier theories of NA functions, GANE emphasizes the effect of NA on gain, which presumably mediates the inverted-U-shaped relation between the efficacy of sensorimotor functions and arousal (Arnsten 2009; Aston-Jones & Cohen 2005). Theories such as network reset and unexpected uncertainty are based on a very distinct intuition: The key role of the LC–NA system in change of internal representations, rather than enhance them (Bouret & Sara 2005; Yu & Dayan 2005). Network reset is based on two features of the NA system: It is extremely well conserved across all vertebrates, and its activation is systematically associated with a profound change in behavior (Bouret & Sara 2004; Clayton et al. 2004; Dalley et al. 2003; Devauges & Sara 1990; Jacobs 1986; McGaughy et al. 2008). The typical condition of LC activation is the orienting response to a salient stimulus (Aston-Jones & Bloom 1981; Bouret & Sara 2004; Foote et al. 1980). Unexpected uncertainty is based on similar intuitions and emphasizes the role of NA in learning (Yu & Dayan 2005). Again, there are some differences between neurobiological intuitions proposed in GANE versus network reset, but the key question the authors raise is not "how," but "why": "Why phasic arousal induced when encountering emotional stimuli can enhance processing of preceding stimuli when they have high priority" (sect. 8.2).

That question implies two features: First, the processing of stimuli is taking enough time to allow subsequent emotional stimuli to induce enhancement of this processing via an increase in arousal. This assumption makes strong predictions on the dynamics of these processes, and indeed, such an assumption is important in understanding LC/NA functions. Second, we are not a priori interested in the original stimulus but the next stimulus, and the preceding stimulus would be ignored. But if the initial stimulus leaves a trace strong enough to be integrated with the emotional one, after the reset, the new "functional network" would underlie the processing of both stimuli. In that case, the representation of the original stimulus would be modified (changed qualitatively), not quantitatively (enhanced or decreased).

Is the influence of arousing events qualitative (network reset) or quantitative (GANE)? Using the example in Figure 7 of Mather et al., these two theories make radically different predictions: According to network reset, the booming sound of a thunderstorm would not enhance the processing of the cow; it would first trigger an orienting response that consists of interrupting existing activity (including processing of the cow) and promoting redirection of attentional resources. Using the words of Mather et al., the sound would become "high priority," but it would either be processed alone or be combined with the cow in a novel representation. Importantly, the representation of the cow as it existed before the storm would disappear.

Thus, we could rephrase Mather et al.'s question: Why should salient stimuli enhance processing of past events? First, time goes one way only, and modulating past events makes sense only if they are used for the present or for planning future actions (James 1913; Sara 2000). The example provided in Figure 7 is very close to laboratory situations in which discrete stimuli are manipulated in a controlled setting. But imagine yourself walking in the fields, and letnow assume that for some reason, you are considering the cow. What will happen if you hear booming thunder? Will you still care about the cow? If yes, what is the chance that you think about it the same way you did before, independently of the critical information provided by the thunder? If the NA system had evolved to enhance the processing of the cow when an inherently more significant stimulus occurs, would this system be so widely represented among living animal species?

In conclusion, in addition to its influence on sensorimotor functions, the LC/NA system has a major role in promoting changes in behavior. The details of the model, including its dynamics, will be critical to understanding how, and why, the release of NA modulates forebrain systems. But this model should account for critical biological features of the LC: It is activated when a behaviorally relevant stimulus triggers a sympathetic response and a behavioral response. For all vertebrates, this autonomic activation is a generic emergency reaction that facilitates coping with a challenge (threat, effort, unexpected event, etc.), and it presumably facilitates the behavioral adjustment to the challenge. This adjustment may take several forms, including gain and/or reset, and be mediated by myriad neurobiological processes, but to understand why the central NA system exists and what it does, it is important to consider ecological problems that the brain has evolved to solve.

The Fluency Amplification Model supports the GANE principle of arousal enhancement
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Abstract: The GANE (glutamate amplifies noradrenergic effects) theory described by Mather et al. offers a neurophysiological basis for the arousal mechanism which is essential for empirical aesthetics and Gestalt processing. More generally, the core principle of perception can be interpreted as a continuous processing of competing arousal states, yielding selective amplification and inhibition of percepts to deduce the meaning of a scene.

The GANE (glutamate amplifies noradrenergic effects) principle Mather et al. describe offers a thorough modeling of how arousal-induced norepinephrine modulates the dynamics of...
information processing. Processing is directed toward high-priority – that is, salient – stimuli, leading to stronger effects on the perception and memory of these stimuli (amplification), whereas the processing of low-priority stimuli is impaired. The recently established fluency amplification model (FAM) (Albrecht & Carbon 2014), originating from the domain of perceptual and affective sciences, builds on a very compatible mechanism. (Cognitive) fluency refers to the experienced ease with which something is processed, mostly operationalized by processing speed or ease of response generation. Typically, theories on fluency assume that the more fluently a stimulus is processed, the higher the appreciation of this stimulus is. In contrast, FAM interprets fluency in terms of saliency: Fluently processed stimuli are more unambiguous and clearer – they are better representatives of their category. As such, the more fluent the processing of a stimulus the stronger the signal is and, thus, the higher the saliency regarding the stimulus is. In FAM, we propose that this causality leads to an amplification of the original judgments assigned to the stimulus. For example, the assessment of the valence of a fluent stimulus will be an amplified version of the initial stimulus valence: Positive stimuli will be valued even more, and negative stimuli will be devalued in a more intense way when being processed fluently. That effect was exactly what we were able to experimentally confirm for stimuli with positive versus negative valence (Albrecht & Carbon 2014). Meanwhile, the emotional assessments of stimuli with minimal saliency, in our case stimuli of undetermined valence, were not altered by fluency. Taken together, these results indicate that the saliency of a stimulus, defined as the deviation from the neutral information regarding the target scale, operates as an amplification factor for the base signal, here, the emotional value of the stimulus.

Beyond FAM, the concept of arousal can also be seen as one essential mechanism underlying amplification effects regarding judgment in general (see, e.g., Storbeck & Clore 2005). The GANE principle offers a plausible neurophysiological basis for a mostly very vaguely defined arousal mechanism that is often used in theories of cognitive sciences. For example, arousal is purported to play an important role in the specific effects of empirical aesthetics (e.g., the misattribution of an internal state – due to unspecified arousal – toward the preference of an object). Arousal is even more influential in the general field of object recognition, where it is assumed to be the key to pooling cognitive resources to increase the probability of solving a perceptual problem, detecting a Gestalt, or recognizing an object. Muth et al. (2015) recently proposed a model explaining the connection between insights (the “aesthetic aha,” which goes along with a sudden rise in fluency [Muth & Carbon 2013]) and preference formation. Aesthetic stimuli are often complex at first sight and difficult to process, which means that they are initially disfluent. Such disfluency indicates the complexity of the perceptual problem, which, in return, signals something potentially meaningful and therefore evokes an orienting reaction plus a state of increased arousal. High arousal shifts attentional and cognitive resources toward the apparent source of complexity, giving rise to interest. Likewise, the GANE model proposes that top-down attention and perceptual features such as contrast and complexity prioritize the processing of certain stimuli over less salient ones. Further elaboration of the stimulus may indeed lead to a decrease in complexity of the visual scene (e.g., by detection of something meaningful or by clear identification of an object), which goes along with a sudden increase in fluency (Albrecht & Carbon 2014). At the same time, GANE proposes that the processing of salient information is amplified, whereas the processing of less salient information is inhibited. These processes finally result in an insight and the dissolving of arousal.

This, on the one hand, has a rewarding quality (see Van de Cruys & Wagemans 2011) independent of the initial stimulus quality; we call this sudden Gestalt-forming event an “aesthetic aha!” or “Gestalt aha!” (Muth et al. 2013); actually such an aha is also paralleled by higher liking of the Gestalt- versus the non-Gestalt-like display (Muth et al. 2013). This rewarding process points to an essential and very general mechanism of perception: to let people continuously seek meaning in visual displays. On the other hand, the rise in fluency facilitates faster and easier processing of the stimulus, resulting in a clearer representation of it, which allows for a more precise, amplified judgment in terms of FAM. Within this scope, the GANE model could help complement the so-called “amplified processing” (Muth et al. 2013) which is interpreted by the GANE model as the intertwining process of selective amplification and inhibition to obtain the most clear interpretation of a given (e.g., visual) scene (Carbon 2014; Gregory 1970) and, thus, to enable the most appropriate action.

Bodily arousal differentially impacts stimulus processing and memory: Norepinephrine in interoception

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Abstract: Bodily arousal modulates stimulus processing and memory, contributing to expression of emotional salience. The “glutamate amplifies noradrenergic effects” (GANE) model proposed by Mather and colleagues can be extended to account for the differential impact of interoceptive (notably cardiac afferent) signals on sensory processing. However, some emotion-specific effects, for example, for fear, may further depend on functional anatomical organisation of affect-related brain structures.

Mather and colleagues provide a compelling account of how stimulus processing is selectively prioritized through interaction of central noradrenaline (norepinephrine) and glutamate release. Their model explains discrepancies regarding the impact of central arousal on aspects of emotion, perception, and cognition. Thus, arousal sometimes enhances the processing of salient stimuli at the expense of neutral or contextual information, while, in other circumstances, it facilitates the processing of neutral stimuli and peripheral information. States of physiological arousal in the body evoked similar psychological effects, suggesting a common mechanism.

In presenting the GANE model, Mather and colleagues refer to studies of skydiving, threat response, processing of emotionally salient (alarming, exciting or disturbing) stimuli, and loud noises. States of running and even unanaesthetized wakefulness in animal experiments are also considered. Arousal is proposed to be a common feature, operationally defined by noradrenaline release from the locus coeruleus. Such states of emotional and behavioural arousal are characterized by physiological changes in the periphery. Within the cardiovascular system, arousal is an embodied action-ready state: Heart rate and blood pressure increases are brought about by enhanced sympathetic drive, parasympathetic withdrawal, and baroreflex inhibition. Bodily arousal feeds back to influence perception, cognition, and emotion, and cardiac and arterial baroreceptors, which fire cyclically on each heartbeat, are a major source of these interoceptive influences.

Relevant to the GANE model, brainstem noradrenergic nuclei including locus coeruleus are sensitive to different interoceptive signals concerning bodily arousal. These nuclei support both...
descending control of autonomic function (A1 and A2 groups within medulla) and ascending control of alertness (e.g., A4 and A6 groups, including nucleus coeruleus). Correspondingly, they react to behavioural challenges by increasing sympathetic drive to the body and by increasing noradrenaline release in the brain via ascending projections from locus coeruleus to hypothalamus, thalamus, and forebrain (cortex and amygdala). Cardiovascular arousal is conveyed to the brainstem in a pulsatile manner by vagus nerve and glossopharyngeal afferents carrying the phasic discharge of baroreceptors that encode the timing and strength of individual heartbeats. The firing of locus coeruleus neurons is regulated by baroreceptor firing (Svensson 1987), resulting in cyclical inhibition of neural activity at late diastole (Elam et al. 1984; 1986; Morilak et al. 1986; Murase et al. 1994). Cardiac afferents modulate activity of nearby brainstem reticular nuclei (Lambertz & Langhorst 1995) and even the amygdala, where the effect is also influenced by state of alertness (Lambertz et al. 1995). Vagus nerve stimulation enhances release of noradrenaline within the amygdala (Hassert et al. 2004). Fine-grained signals concerning bodily arousal can thus influence perception and cognition via brain regions governing alertness and central arousal. Baroreceptor signals occurring with each heartbeat impact stimulus detection (Garfinkel et al. 2014; Park et al. 2014), memory (Garfinkel et al. 2013), and emotional responses (Garfinkel et al. 2014). Yet when it comes to processing emotional information, these physiological arousal signals evoke selective effects. Although cardiac systole inhibits the processing of pain stimuli (Gray et al. 2009) and attenuates the encoding into memory of words irrespective of valence (Garfinkel et al. 2013), the processing of fear stimuli is enhanced (Garfinkel et al. 2014).

The emotional attentional blink paradigm illustrates the prioritised processing of emotional stimuli. At the limit of perceptual awareness, emotional stimuli can overcome a perceptual block, the attentional blink effect, breaking through to awareness by capturing attention. This index of emotional salience is adrenergically mediated, being enhanced by administration of the noradrenergic reuptake inhibitor reboxetine and abolished by ß-adrenoceptor blockade with propranolol (De Martino et al. 2008). This prioritised processing of emotional stimuli also depends on the functional integrity of the amygdala (Anderson & Phelps 2001). The additional impact of afferent signals concerning cardiovascular arousal on early affective processing can be measured by timing the presentation of target stimuli to distinct phases (systole and diastole) of the cardiac cycle. Here the outstanding observation is a selective cardiac enhancement of fear processing, manifest in the emotional attentional blink task as better detection of fearful faces presented at systole, compared with diastole. This cardiac cycle effect is not seen for disgusted, happy, or neutral faces (although there is a trend for neutral faces to be better detected at diastole) (Fig. 1) (Garfinkel et al. 2014). Moreover, at systole, increased amygdala activity in response to fear compared with neutral stimuli predicts increased subjective rating of fear intensity and underscores the selective contribution of cardiac afferent signals to amygdala-mediated processing of salient stimuli (Garfinkel et al. 2014).

Thus, interoceptive signals concerning cardiovascular arousal can both increase (e.g., fear) and decrease (e.g., words, pain) stimulus processing. This is differentiated by the type of task or the emotion class of the stimulus. Although the GANE model explains much of the differential impact of cardiac afferent signals on sensory processing, it only partially accounts for emotion specificity and (task-related) behavioural demand that can further differentiate and guide the directionality of arousal effects. Encompassing physiological state within the concept of arousal reveals levels of interaction and a selective impact of the arousal signal itself. The glutamate component of the GANE model takes into account prioritisation of certain stimulus types, yet it underplays the degree to which this specificity must also depend on the differential anatomical organisation of critical brain structures supporting emotion-related response repertoires.

Amplified selectivity in cognitive processing implements the neural gain model of norepinephrine function
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Abstract: Previous work has suggested that an interaction between local selective (e.g., glutamatergic) excitation and global gain modulation (via norepinephrine) amplifies selectivity in information processing. Mather et al. extend this existing theory by suggesting that localized gain modulation may further mediate this effect—an interesting prospect that invites new theoretical and experimental work.

Mather and colleagues’ article joins the growing body of work suggesting that norepinephrine, through its brainwide effect on neural gain, selectively enhances useful and salient neural representations (Aston-Jones & Cohen 2005; Eldar et al. 2013; Usher et al. 1999). Building on an early computational model of catecholamine function (Servan-Schreiber et al. 1990), and later work directly addressing locus coeruleus function (Usher et al. 1999), Aston-Jones and Cohen (2005) proposed that one of the roles of the locus coeruleus–norepinephrine system is to enhance, through gain modulation, neuronal representations that are most useful for maximizing utility (adaptive gain theory).
Critically, although norepinephrine is released globally throughout the brain, it was argued that its effects could be temporally and spatially specific. Temporally specific because norepinephrine can be phasically released in response to task-relevant stimuli and, thus, suitably timed to enhance representations that are most useful for task performance. Spatially specific because gain modulation inherently entails an interaction between norepinephrine and glutamate in which strong neural representations (i.e., those that are already receiving strong glutamatergic input, because of “bottom-up” sensory inputs and/or “top-down” context or control) are enhanced by norepinephrine, whereas weak neural representations are more inhibited (Eldar 2014; Eldar et al. 2013; see also Figure 5 in Mather et al.).

We conducted a series of behavioral and neuroimaging experiments to test this idea, that norepinephrine amplifies selectivity in information processing (Eldar 2014). Specifically, we investigated the relationship between selectivity and pupilometric indices of norepinephrine function in the domains of learning, perception, and memory. We first showed that indices of high norepinephrine function are associated with learning that is more selectively focused on stimulus features to which individuals are predisposed to attend (Eldar et al. 2013). We then showed that a similar effect is evident in the domain of perception. Specifically, we found that indices of high norepinephrine function are associated with perception of ambiguous characters that is more selectively focused either on the character’s visual features or on its semantic context, depending on which source of information has stronger influence (we manipulated the source’s strength using subliminal priming [Eldar 2014; Eldar et al., in press]). Notably, the latter finding suggests that norepinephrine will enhance bottom-up (e.g., visual features) or top-down (e.g., semantic) influences on perception, whichever is stronger. Finally, we also showed that a similar effect is evident in the domain of memory, where we found that indices of high norepinephrine function are associated with recognition memory that is more selective to the font in which a word appears, when attention is drawn to the font by the experimental task (Eldar 2014; Eldar et al. in press). These findings of increased selectivity in learning, perception, and memory were predicted by neural network models of norepinephrine function in which the effect of norepinephrine was modeled as a global increase in gain.

In addition to the behavioral predictions, our neural network models generated several neural predictions, which we tested using functional magnetic resonance imaging. First, increased gain entails that neural activity should be driven to maximal and minimal levels, and thus, the absolute deviation of activity levels from mean activity should increase with gain. Second, stronger responsiveness to input signals should increase functional connectivity between neural units. Third, functional connectivity between neural units should become more selectively localized in clusters (i.e., less globally distributed), mirroring the behavioral selectivity that is associated with high gain. Indeed, pupilary indices of high norepinephrine function were associated with all three effects throughout the brain, as measured by brainwide blood oxygen level-dependent (BOLD) signals, further supporting the role of norepinephrine in global gain modulation in humans (Eldar et al. 2013).

The gain modulation model of norepinephrine function was originally inspired by findings that norepinephrine enhances single-neuron responses to both excitatory and inhibitory signals (e.g., Moises et al. 1979; Waterhouse & Woodward 1980), which suggested that norepinephrine increases the contrast between strongly and weakly active neurons. However, subsequent single-neuron electrophysiology studies showed that norepinephrine may either enhance or suppress responsiveness to excitatory input, depending on which receptor it activates (e.g., Devilbiss & Waterhouse 2000). Mather and colleagues’ proposal of local positive-feedback interaction between norepinephrine and glutamate reconciles this latter evidence with the neural gain model of norepinephrine function, because it suggests a mechanism through which the gain-enhancing effect of norepinephrine would dominate specifically in strongly activated neurons, and thus, norepinephrine’s overall effect would be to increase the contrast between weakly and strongly active neurons, as in the original model (shown in Fig. 5 in Mather et al.). In addition, the local changes in norepinephrine that Mather et al. propose may have additional effects that go beyond those of the interaction between local excitation and global gain modulation. For instance, local enhancement of gain may amplify selectivity even further. Indeed, such local changes have been suggested by early in vivo studies of the influence of sensory and thalamic inputs on cortical release of norepinephrine (e.g., Marrocco et al. 1987).

In sum, the neural gain model of norepinephrine function has been successful in predicting a range of norepinephrine’s neural and behavioral effects, among which is amplified selectivity in perception and memory. Mather and colleagues’ proposal of local glutamate–norepinephrine interaction further supports the neural gain model, suggesting that additional local interactions may enhance this effect. This suggestion invites further modeling to generate quantitative predictions and experimental work to test them.

The role of arousal in predictive coding

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Abstract: Within a predictive coding approach, the arousal/norepinephrine effects described by the GANE (glutamate amplifies noradrenergic effects) model seem to modulate the precision attributed to prediction errors, favoring the selective updating of predictive models with larger prediction errors. However, to explain how arousal effects are triggered, it is likely that different kinds of prediction errors (including interoceptive/affective) need to be considered.

Classical models of information flow in the cerebral cortex consider that primary sensory regions detect the physical properties of the stimuli which are then combined into increasingly complex representations along the hierarchy of perceptual processing. As such, on the one hand perceptual processing is considered to be largely bottom-up, and top-down effects are expected to modulate the processing stream only. On the other hand, the predictive coding framework suggests that the cortical representation of objects is produced largely by top-down feedback to sensory cortices (i.e., predictions about what is being perceived originate in higher-level regions) (Clark 2013; Friston 2005; 2010). In this view, sensory information is not fed forward along the cortex, but rather, what is communicated along the cortical hierarchy is only the difference between the predicted and actual inputs: the prediction errors. When such a mismatch occurs, the prediction errors are then used to update the higher levels of the hierarchy to update the predictive model so as to eliminate prediction errors in the next round of comparisons (Clark 2013; Huang & Rao 2011; Rao & Ballard 1999). Predictions and prediction errors are thought to be instantiated by different neural units, and the balance between the two depends on precision cells that modulate their relative weights. Increased precision of the prediction errors means that the error signal will be strengthened by the precision units and lead to a stronger updating of the predictive model, whereas decreased precision suppresses the prediction errors and, thus, maintains the current model (Barrett & Simmons 2015). With this brief introduction in mind, I now turn to how the GANE (glutamate amplifies noradrenergic effects) model may be integrated within a predictive coding approach – a possibility that is acknowledged by Mather et al.

The activity of norepinephrine (NE) neurons has been in the focus of researchers interested in the neural coding of prediction...
errors (Dayan & Yu 2006). NE cells respond phasically to unexpected stimuli across sensory modalities and cease to respond after a few repetitions of the stimulus, a pattern of activity that is consistent with what would be expected from units coding prediction errors (Schultz & Dickinson 2000). As detailed by the GANE model, however, the overall effect of NE seems to be more akin to a modulation of the precision weights of prediction errors. Thus, in predictive coding terminology, NE amplifies the stronger feedforward glutamatergic error signals while suppressing weaker prediction errors, leading to a stronger updating of only the most unexpected inputs. Indeed, this is a sensible explanation: strong prediction errors signal highly unexpected sensory input and, thus, elicit orienting responses and concomitant central NE release to boost signal-to-noise ratio and favor the updating of the most relevant predictions.

The salience or priority of stimuli that seem to trigger NE effects, however, is not fully dependent on sensory mismatch. It is true that phasic NE responses occur to intense unexpected sensory inputs (Petersen & Posner 2012), but also to stimuli that are not physically extreme, namely, stimuli that carry emotional or task-related significance (Schulz & Dickinson 2000). Indeed, the affective/motivational aspect of arousal is something that has not been the focus of the more classic formulations of predictive coding approaches. However, recent models of affective predictive coding extend the predictive coding framework, originally developed to account for perception of external objects, to include interoception, that is, the cortical representation of internal states that constitute the basis of emotional experience (Barrett & Simmons 2015; Seth 2013). Also, affective predictive coding models do not consider interoceptive inferences as independent from exteroceptive processing, but rather consider that affective predictions and affective prediction errors are basic components of “regular” perception (Barrett & Bar 2009). This means that the perception of an object involves predictions not only about its physical features (e.g., shape, color), but also about its affective properties (e.g., very pleasant, neutral, scary), and that the prediction errors that are elicited may concern sensory and affective mismatches.

One hypothesis consistent with this view is that engagement of NE neurons in the locus coeruleus may depend on a threshold of the net sum of prediction errors for a given input. This would mean that arousal effects may occur following sensory, affective, or task-related mismatch (depending on whether the stimulus is, respectively, inconsistent with perceptual, interoceptive, or goal-related predictions) or a combination of these. If this combination of prediction errors reaches a given threshold, then a phasic NE response is elicited to facilitate the selective updating of predictions in the prioritized manner that Mather and colleagues elegantly describe. Indeed, it has been reported that emotionally deviant stimuli evoke larger cortical prediction errors than neutral deviants (Vogel et al. 2015a), but the precise role of NE in this effect remains an issue for future investigation.

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Are there “local hotspots”? When concepts of cognitive psychology do not fit with physiological results

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Abstract: Mather and colleagues’ arguments require rethinking at the mechanistic level. The arguments on the physiological effects of norepinephrine at the cortical level are inconsistent with large parts of the literature. There is no evidence that norepinephrine induces local “hotspots”: Norepinephrine mainly decreases evoked responses; facilitating effects are rare and not localized. More generally, the idea that perception benefits from “local hotspots” is hardly compatible with the fact that neural representations involve largely distributed activation of cortical and subcortical networks.

Mather and colleagues propose that phasic activation of LC neurons biases perception and memory. They suggest that elevated levels of glutamate at the site of prioritized representations increase local norepinephrine (NE) release, creating “NE hotspots.” At these spots, enhancement of glutamate and release of NE mutually enhance and amplify the activation of prioritized representations. This excitatory effect contrasts with the widespread suppression by NE of weaker representations via lateral and auto-inhibitory processes.

Mather et al. provide a schematic representation at the cellular level (Fig. 6), but is it supported by physiological data obtained in sensory cortices? Although the locus coeruleus (LC) neurons project widely to many cortical areas, recent data indicate that some neurons project more to one area (the prefrontal or motor cortex) than others (Chandler et al. 2013; 2014). Within an area, NE is released in the extracellular space from NE varicosities and reaches the entire cortical network. When sensory stimuli are processed by cortical neurons, glutamate is released by the thalamocortical terminals. How does NE affect cortical processing? Iontophoretic application of NE performed in the somatosensory, visual, and auditory cortices revealed that, in most of the cases, NE depressed evoked responses (e.g., Kolta et al. 1987; Manunta & Edeline 1997; 1998; Videen et al. 1984), an effect replicated in awake animals (Bassant et al. 1990; Foote et al. 1975; Manunta & Edeline 1999). Moreover, in awake rats, tonic activation of LC neurons by continuous low-frequency stimulation (1 Hz) triggered similar effects: decreased evoked responses in 63% of cells in the rat somatosensory cortex (Devillibus & Waterhouse 2004). It has been argued that these inhibitory effects were a consequence of the very high concentrations of NE in the vicinity of the cell (Waterhouse et al. 1998a), but this seems unlikely given that pronounced depression of evoked responses was also observed with very low ejection currents (Ego-Stengel et al. 2002; Manunta & Edeline 1997; reviewed in Edeline 2012). If the hotspot theory were the main mechanism at play, then exogenous application of NA would more likely have increased evoked activity.

But what are the consequences for the neurons’ functional properties? In the auditory cortex, the suppressive effect of NE promotes an increase in frequency selectivity in both anesthetized and unanesthetized animals (Edeline 1995; Manunta & Edeline 1997; 1999). In the visual cortex, application of NE improved the velocity and direction selectivity of cells, without modifying orientation selectivity (Ego-Stengel et al. 2002; McLean & Waterhouse 1994). These results clearly point out that the effects of NE (and other neuromodulators) can differ depending on the stimulus dimension. For example, a dimension that depends on thalamocortical affinances (such as frequency tuning in the auditory cortex or size of the receptive field in the visual cortex) could be more affected than a dimension that relies more on corticocortical affinances (such as frequency modulation tuning in the auditory cortex or velocity tuning in the visual cortex). Yet, glutamate is released in all cases, indicating that the glutamate–norepinephrine interaction is not as straightforward as described by the authors.

One may ask if it is possible for “NE hotspots” to emerge when NE is repeatedly associated with glutamate release at particular synapses? When a stimulus that activates a specific set of synapses is paired with phasic LC stimulation, a predominant decrease in neuronal activity is initially reported in several cortical areas (e.g., Olpe et al. 1980; Sato et al. 1989). In the somatosensory cortex, both the excitatory and inhibitory components of evoked responses are facilitated when phasic stimulation of the LC is
Contemplating the GANE model using an extreme case paradigm

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Abstract: Early experiences play a crucial role in programming brain function, affecting selective attention, learning, and memory. Infant http://www.cambridge.org/core/terms. https://doi.org/10.1017/S0140525X15001843

literature suggests an extension of the GANE (glutamate amplifies noradrenergic effects) model to conditions with minimal priority-map inputs, yet suggests qualifications by noting that its efficacy is increased when tonic levels of arousal are maintained in an optimal range, in manners that are age and exposure dependent.

Mather and colleagues' intriguing GANE (glutamate amplifies noradrenergic effects) model underscores an important process, through which GANE changes influence the selection process to favor high- over low-priority representations.

The extended literature covered in the article concentrates mostly on experimental research with typically developing young adults, whose performance relies on an established neural network, set with implicit “know-hows” and an explicit knowledge base, which shape and set local hotspots, to be activated proactively in the prospect of newly arriving inputs (Bouret & Richmond 2015). One way to test this model may be an extreme case paradigm in which top-down priorities are negligible, and the roles of global brain activation are augmented, as is the case of the newborn.

Research on infancy, early development of attention, and arousal, in typical and clinical samples opens the discussion of the generalizability of the GANE model because young infants perceive stimuli with no preset priorities and with little previous knowledge. As such, infancy offers an interesting test case for the GANE model.

Early-life experiences play a crucial role in programming brain function, particularly with respect to selective attention, learning, and memory (Geva et al. 2006). Newborns are busy perceiving and memorizing the environment at rates that are not surpassed thereafter, equipped with an impressionable template that does not allow yet for exerting deliberate priority operations. How might GANE function at infancy? Models with neonates highlight four interdependent notions that may qualit the limits of the proposed model, with respect to development, exposure, global activity, and resilience to variance.

Dependence on development. Neonates and adults differ markedly in their ability to learn selectively (Kuhl et al. 1992). These differences were suggested to be related, in part, to developmental differences in arousal response to sensory stimuli (Kuhl 2007) as a function of differential locus coeruleus–norepinephrine (LC–NE) activity (Moriceau & Sullivan 2004; Nakamura & Sakaguchi 1990). Differences are such that compared with the LC of the infant, the adult LC gradually becomes less likely to respond to non-noxious stimuli (Kimura & Nakamura 1985; Nakamura et al. 1987; Selden et al. 1990), habituates earlier in response to repeated (or even single) stimulation (Vankov et al. 1995), and produces shortened LC responses in response to sensory stimulation (Nakamura & Sakaguchi 1990). All of these differences suggest a potential role for development in the proposed model (Moriceau & Sullivan 2004).

Dependence on experience. Exposure at sensitive periods seems to play a significant role in the development of the LC–NE system (Nakamura et al. 1987; Rangel & Leon 1995). Also, experience early in development has been found to affect PFC responsiveness to LC–NE. For example, neonatal experience involving maternal contact reward was reported to affect the noradrenergic system of the rat prefrontal cortex (Kalpachidou et al. 2013). The experience was related to hypermethylation of the β-adrenergic receptor gene promoter and consequently enhanced expression of its mRNA in the prefrontal cortex, resulting in better discrimination and improved learning in the young pups (Kalpachidou et al. 2015).

In addition, selective recognition of maternal odors has been found to be accompanied by increased release of glutamate and GABA from the dendrodendritic synapses and an increased efficacy of glutamate-evoked GABA release (Kendrick et al. 1992), and early-life stress related to maternal separation has been reported to alter glutamate and GABA transmission and, in particular, to alter GABAA receptor expression (Sterley et al. 2013).

Commentary/Mather et al.: Norepinephrine ignites local hotspots of neuronal excitation
The integration of these findings points to the possible role of early-life exposure in the GANE model. **Dependence on tonic levels of activity.** The LC is thought to play a central role in regulating arousal states in addition to its role in attention and memory (Howells et al. 2010; Rajkowski et al. 1994). Initial leads from human infancy research point to the notion that in the case of the newborn, arousal homeostasis possibly plays a significant role in attention and in recognition (Geva et al. 1999), with brainstem pathways playing a central role in gating arousal self-regulation (Geva & Feldman 2008). Feeding-dependent arousal differences were found to affect newborn preferences for cognitively demanding stimuli (Geva et al. 1999; 2013), the interaction is such that more aroused neonates tend to orient toward less intense familiar stimuli; yet when less aroused, newborns prefer more intense stimuli (Gardner & Karmel 1983; 1984) and orient toward novel stimuli as compared with familiar ones in visual recognition memory tasks (Geva et al. 1999).

Sleep-wake arousals also seem to play a similar role. Recent work with intracellular recordings has shown an interaction of LC activity in monkeys as a function of fatigue, an effect attributed to the LC possibly providing the impetus to act when the predicted outcome value is low (Bouret & Richmond 2015). Indeed, arousal states were found to affect attention in young human infants. Neonatal sleep fragmentation was reported to be associated with infants’ focused attention to specific stimuli early in development (Geva et al. 2013). Compared with good sleepers, infants who were poor sleepers as neonates had difficulties focusing on target stimuli in the presence of complex distracters, but managed focusing in the presence of simpler distracters. Integration of the findings on arousal state effects on attention and memory emphasizes the need to consider tonic arousal changes in the GANE model.

Finally, the validity of the model may gain from testing of its limits in neuropsychiatric disorders, such as attention-deficit/ hyperactivity disorder (Sterley et al. 2013), which involve poor adaptation to change (Sara 2009). Such an exploration may suggest the notion that GANE efficacy is increased when tonic levels of arousal are maintained in an optimal range.

Together, these data suggest an extension of the GANE model to infancy; however, integration of the above findings with the framework presented suggests a qualification to the GANE model, by noting that its efficacy is increased when tonic levels of arousal are maintained in an optimal range, in manners that are age and exposure dependent.

**Dentate gyrus and hilar region revisited**

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Abstract: It is suggested that the dentate gyrus and hilar region in the hippocampus perform memory selection and that the selectivity of the gating of memory by this circuit is modulated by the norepinephrine–glutamate loop described by Mather et al.

Mather et al. propose that arousal modulates attention through a norepinephrine–glutamate feedback loop in local circuits. Here, I suggest a specific circuit where this mechanism may be in operation: the granule cell–mossy cell loop in the hippocampus.

It is commonly proposed that the CA3 region of hippocampus forms an associative-memory store for short- and medium-term memories (Gardner-Medwin 1976; Hopfield 1982; Levy & Steward 1979; Marr 1971; McNaughton & Morris 1987; Rolls 1989; Treves & Rolls 1992). Here, pictures, memories, in the form of patterns of activity in the entorhinal cortex, feed forward along the perforant pathway to CA3, activating a sparse subset of the CA3 pyramidal cells. Plasticity in the synapses of the recurrent network in CA3 and in the perforant pathway synapses onto CA3 neurons fixes the memory so that it can be recalled. If a part of the same pattern of activity occurs in entorhinal cortex, the corresponding part pattern is activated in CA3 and it is then completed by auto-associative dynamics.

Pattern collision, where two similar memories are confused during pattern completion, is a problem in auto-associative networks, particularly if they are required to rapidly store memories with only a small number of presentations. It is likely that the hippocampus has a mechanism to avoid or reduce pattern collision: the hippocampus stores rapidly acquired memories, and it is important that similar but distinct memories can be distinguished during recall.

It has been proposed that the role of the dentate gyrus is to separate patterns and thereby reduce collisions (Gilbert et al. 2001; Leutgeb et al. 2007; McHugh et al. 2007; O’Reilly & McClelland 1994; Treves & Rolls 1992). In addition to CA3 neurons, the perforant pathway connects to the granule layer in dentate gyrus. The LC is thought to have a key role in governing the pattern of differences in the LC, in turn, connected to CA3 along the mossy fibers. This means that the entorhinal cortex is connected to CA3 directly, along the perforant pathway, and indirectly, via dentate gyrus. In the specific case of dentate gyrus pattern separation proposed by O’Reilly and McClelland (1994), there is local k-winner-take-all dynamics between cells in dentate gyrus, and the consequence of this is that only a random subset of the cells receiving input from entorhinal cortex become active. This activity is fed forward along the mossy fibers to CA3 and, in turn, excites a random subset of those cells in CA3 that receive input from entorhinal cortex. This randomization separates the patterns that are then learned in the CA3 auto-associative network.

There is experimental evidence (McHugh et al. 2007) that the dentate gyrus is important for pattern separation and that the adult neurogenesis of dentate gyrus granule cells, which may support the randomization, is linked to pattern separation (Altman 1963; Bayer et al. 1982; Clelland et al. 2009; Salbay et al. 2011). However, it seems unlikely that pattern separation is the only role of the dentate gyrus; for a start, pattern separation on its own seems a modest role for such a substantial brain region. Beyond this, pattern separation does not explain either the hilar region or the role of norepinephrine in the dentate gyrus.

The hilar region lies between dentate gyrus and CA3. As the mossy fibers run through the hilar region they form en passant connections with the mossy cells (Amaral 1978; Scharfman & Myers 2013). These are large excitatory cells whose proximal dendrites are covered in mossy-looking spines. The mossy cells, in turn, have a substantial backprojection that extends along the longitudinal axis of the dentate gyrus and CA3 (Altman & Witter 1988; Amaral et al. 2007) and connects to both granule cells and inhibitory interneurons (Scharfman 1984; 1995).

This two-layer structure seems more elaborate than a simple randomizing k-winner-takes-all network would require; random subselection from a pattern could be achieved by local excitatory–inhibitory dynamics within the dentate gyrus itself. However, the two-layer structure would make sense if the role of the dentate gyrus encompassed memory selection as well as pattern separation. As pointed out by Koch et al. (Koch & Ullman 1984; 1987; Olshausen et al. 1993), a single layer winner-takes-all network in which competition occurs across the whole network requires considerable interneuronal connectivity. This issue is resolved by having more than one layer; in the first layer, competition is restricted to subregions, and a champion emerges from each subregion to compete in the next layer where the competition between subregions occurs. In short, I suggest here that, in addition to separating patterns, the winner-take-all dynamics in the dentate gyrus also compares the salience
of different aspects of its input and that this selection gate refines the storage of memories in CA3. The role of the hilar region is to facilitate this comparison.

The locus coeruleus projects to the dentate gyrus, which contains β-adrenergic receptors (Berridge & Waterhouse 2003; Harley 2007). Norepinephrine release in response to novelty during exploration enhances excitability in the dentate gyrus (Kitachigina et al. 1997); in fact, the activity of both interneurons (Nitz & McNaughton 2004) and excitatory neurons (Dahl & Winson 1985; Neuman & Harley 1983) in dentate gyrus show norepinephrine-promoted increase in response to novelty. Furthermore, it has been reported that in hippocampus, glutamate causes enhanced norepinephrine release (Pittaluga & Raiteri 1990; Raiteri et al. 1992), an effect that is most marked in the dentate gyrus (Andrés et al. 1993). Conversely, norepinephrine in dentate gyrus, but not in other hippocampal regions, potentiates the release of glutamate (Lynch & Bliss 1986). The role of norepinephrine in dentate gyrus seems somewhat mysterious if the role of the dentate gyrus is restricted to pattern separation. However, if, as proposed here, the dentate gyrus also performs memory selection, then the norepinephrine–glutamate mechanism for modulating memory selectivity described by Mather et al. becomes the missing clue that could explain the role of norepinephrine in dentate gyrus.

**GANEing on emotion and emotion regulation**

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Abstract: The function of emotion and its underlying neural mechanisms are often left underspecified. I extend the GANE (glutamate amplifies noradrenergic effects) model by examining its success in accounting for findings in emotion regulation. I also identify points of alignment with construction models of emotion and with the hypothesis that emotion states function to push neural activity toward rapid and efficient action.

What is emotion? Why did it evolve and what is its purpose? Several models of the origin and function of emotion have been put forward (see Gross & Barrett 2011 for review). For the sake of brevity, I identify two broad and encompassing approaches here. One pursues questions related to why and how emotions evolved (Ekman 1993; Tooby & Cosmides 1990), referred to as the entity view. The other focuses on why emotion evolved (Lindquist et al. 2012; Simón 1967), and that one I will call the process view. At first glance it is a subtle difference, but it is a difference that matters greatly for how we understand emotion, its underlying mechanisms, and its adaptive functions.

From the entity view, individual emotions served specific functions in the past that were important for survival and so were preserved. Each emotion is structured like an organ that has feature detectors for identifying relevant stimuli that then trigger a coordinated set of action tendencies that enhance survival (Panksepp 2007). The challenge for modern humans is to regulate these inherited responses to conform to the much changed present-day environment. The process view, on the other hand, allows for a nearly unlimited variety of emotional states and responses. Principles of neural computation are often an important part of this account (Lindquist et al. 2012), and can be augmented by stipulating that the function of emotion is to enable neurologic systems to minimize exploration over the current problem space to more rapidly bring about efficient action (cf. Donoso et al. 2014; Simón 1967), something we refer to as computational expedience.

The GANE model is an intriguing mechanism, supportive and suggestive of the process view of emotion insofar as it binds any variety of prioritized cortical representations to arousal and core affective states, rather than assuming that an individual category of emotion (sadness or fear, for example) produces stereotyped cognitive and behavioral effects. Thus, GANE suggests that there is a great deal of flexibility in the formation of emotion states and that no special neural substrates or modules of particular emotions are needed to account for the adaptive, and sometimes maladaptive, nature of emotional memory and regulation. The process view is consistent with the increasingly influential models of emotion construction; however, a full exposition is beyond the scope of this commentary.

I instead focus on the principle of computational expendiency within the framework of GANE by drawing on findings from emotion regulation. An inability to modulate arousal may lead to difficulties in adapting to present circumstances, not because emotional states are geared toward environments from our phylogenetic past, but because alternative and more adaptive forms of responding may not reach priority in the cortex. If one’s affective learning history prioritizes maladaptive cortical representations because of social modeling, maltreatment, or potentially traumatic low-probability events, GANE suggests that unless there is a dampening of arousal, or assistance in pushing subthreshold representations into greater excitement, or both, it will be difficult to alter behavioral responses. Less clinically, because the neural categorization of stimuli and situations is probabilistic (Donoso et al. 2014), cortical activity that represents situations will inevitably err from time to time. Giving oneself space to explore alternatives through arousal regulation efforts is likely to help individuals recover from misattributions and behave more adaptively. Reducing arousal through emotion regulation would provide an opportunity for neural activity representing alternatives to reach priority and, thus, have an impact on action and memory.

For these reasons, emotion regulation is a ubiquitous human activity (Gross 2015), and recent research indicates that it can take on a variety of forms depending on the situation and one’s goals. Humans often increase emotional states that they believe will enhance their performance (Tamir et al. 2015). For example, when preparing for an upcoming negotiation, individuals will increase negative emotional states (such as anger) to increase the likelihood of obtaining their goal. We could speculate that the arousal and prioritized representations we label as anger in this instance provide a singularity of focus and purpose unenumerated by the deliberation of alternative states as suggested by GANE mechanisms. Relatedly, the strategy of distraction is more effective for high-intensity stimuli, whereas altering one’s interpretations (a strategy called reappraisal) is more effective for low-intensity stimuli (Sheppes et al. 2014). We suggest that distraction during high-arousal events helps to reduce the tendency toward computational expendiency of prioritized representations, to allow for further exploration of the dangers, demands, and opportunities of the situation. When arousal is low, reappraisal is more successful at prioritizing new representations so that new memories can be formed that change how one would respond to the stimulus in the future (Denny et al. 2015). Importantly, the success of reappraisal is severely impacted if there is high arousal (Raio et al. 2013), perhaps because cortical activity representing alternative interpretations is unable to reach the high-arousal threshold and achieve priority.

A final intriguing case is the impact that cognitive load has on reducing hedonic arousal and temptation (Van Dillen et al. 2013). As Mather et al. note, many models of cognition and emotion assume that emotionally evocative stimuli always take priority over attention. However, findings that cognitive load interferes with the introduction of new emotion states further support the GANE model by demonstrating how current prioritized representations are maintained by arousal to the exclusion of alternative representations, even those that would otherwise be emotional given one’s affective learning history.

It is not my intention to argue that all aspects of emotion, emotion regulation, or psychopathology can be accounted for by the GANE model. Numerous other neurotransmitters and neurons are likely to play important roles as well.
neuromodulators will also play decisive roles. However, the GANE model offers a neural mechanism that helps to unify cognition and emotion while drawing attention to neurocomputational effects that align with previous theorizing on the function of emotion in ways that are suggestive of future research on the mechanistic bases of emotion regulation.

Once more with feeling: On the explanatory limits of the GANE model and the missing role of subjective experience

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Abstract: We applaud Mather and colleagues’ model, which emphasizes the neurobiological pathways by which affective arousal tunes attention and memory. This commentary offers a friendly discussion of several potential limitations of the theory. We suggest the model is strong when predicting task-driven demands but is limited when predicting the impact of individual biases, interpretations, and experiential feelings.

Mather et al. introduce an impressively broad neurobiological model of the role of affective arousal in directing attention and memory. Rather than discuss the many strengths of the GANE model, our commentary offers a friendly discussion of some limitations of the model and the missing role of subjective experience.

One concern is the predictive utility of the model. The model can account for a variety of effects, but it fails at making clear a priori predictions for attention and memory effects. One reason it lacks predictive utility is its reliance on salience. The use of salience falls victim to a circular argument, because salience often depends on confirmation from the results (self-dependent justifications) (Hahn 2011). Such circularity hampers theory prediction because salience concedes vagueness as it becomes defined post hoc or through task demands. For instance, imagine that participants were asked to attend to a central fear face and ignore surrounding faces. One study used neutral faces as distractors and found better attention/memory for fear faces, but another study used angry faces as distractors and found better attention/memory for angry faces. The vagueness of saliency allows for both studies to support the model (saliency determined by task demands and stimuli, respectively); yet, a naive researcher would fail to make these distinct predictions with the GANE model. Thus, the model can account for various effects, but fails to make clear, deductive predictions (such circularity plagued the depth of the processing approach) (Craik & Lockhart 1972).

The model also does not address predictions based on individual differences. If we compared memory for task-dependent salient stimuli (snakes) in the face of distractors (spiders), at a group level, people may show better memory for snakes than spiders. However, would this be true for each individual? Probably not. Spider-phobics may remember the spider rather than the snake. Therefore, can the model accurately predict when task demands or individual biases will have a greater impact on attention/memory? Moreover, can this model be extended to predict attentional/memory biases linked to various mental health disorders (depression, anxiety, attention deficit/hyperactivity disorder)?

Attention and memory are treated objectively in the model, but they are in the spirit of improving the GANE model. Indeed, there is much to like about the model and we agree more than disagree with much of it.
Interactions of noradrenaline and cortisol and the induction of indelible memories

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Abstract: The glutamate amplifies noradrenergic effects (GANE) model emphasizes the role of local glutamate–noradrenergic interactions in creating functional hotspots for prioritized processing of salient stimuli. Here, we briefly outline current evidence that synergistic action of noradrenaline and cortisol enables emotional stimuli to gain privileged access to amygdala–hippocampus circuits, eventually resulting in the formation of indelible memories and posttraumatic stress disorder (PTSD).

In their superb glutamate amplifies noradrenergic effects (GANE) model, Mather and colleagues convincingly argue that under conditions of arousal-induced phasic activity of the locus coeruleus (LC), locally elevated glutamate (GLU) levels amplify noradrenergic (norepinephrine [NE]) release from the LC, thus creating functional hotspots of prioritized processing that bias perception and memory. Although the GANE model focuses on stimulus salience coding through rapid GLU and NE signaling and their focal interactions, it should be emphasized that endocrine signals, including the adrenal stress hormone cortisol (CORT), brain concentrations of which peak within minutes as a result of hypothalamus–pituitary–adrenal (HPA) axis activation (de Kloet et al. 2005), also intimately interact with NE to code perceptual and mnemonic priority, especially under conditions of emotional arousal.

In functional magnetic resonance imaging (fMRI) experiments, emotional arousal is frequently operationalized by exposing subjects to facial displays of emotion, which evoke responses in specific functional subdivisions of the amygdala (Goossens et al. 2009; Hurlemann et al. 2008). One established means of segregating the neuromodulatory effects produced by NE, CORT, and their interactions, is pharmacologic fMRI (phMRI) (Patin & Hurlemann 2011). A combination of phMRI with histoprobabilistic maps of the subregional architecture of the amygdala (Goossens et al. 2009; Hurlemann et al. 2008) revealed that blockade of β-noradrenergic receptors with the non-specific antagonist propranolol (40 mg po) desensitized the basolateral amygdala (BLA) (Hurlemann et al. 2010), which is consistent with behavioral data indicating that propranolol (40 mg po) eliminated a facilitation of declarative learning from facial feedback (Mihov et al. 2010). In contrast, enhancement of BLA reactivity with the NE re-uptake inhibitor (NARI) reboxetine (4 mg po) produced a response bias toward fearful faces (Onur et al. 2009). Together, these results suggest that increases in NE signaling may be essential for converting the BLA—an area of the brain controlled by powerful inhibitory circuits (Ehrlich et al. 2009)—into a fear module (Onur et al. 2009). One interpretation of these findings is that phasic increases in endogenous NE signaling per se might be sufficient to code stimulus salience. However, because of its pivotal role in orchestrating fear memory acquisition and storage via N-methyl-D-aspartate (NMDA) receptor-mediated long-term potentiation (LTP) (Ehrlich et al. 2009), the BLA may be a locus of extensive GLU–NE interactions, such that observations of a reboxetine-induced increase in BLA signals may, in fact, support the GANE model.

In addition to rapid neuromodulatory effects mediated by NE per se, emotional arousal elicits heightened adrenal release of CORT, which feeds back on the amygdala and hippocampus via activation of mineralocorticoid and glucocorticoid receptors in these regions (de Kloet et al. 2005; McEwen et al. 2015). Experimentally, this endocrine response can be mimicked by exogenous administration of synthetic CORT (20–40 mg po), and studies based on this challenge not only have noted a desensitization of the amygdala during fear conditioning (Merz et al. 2010) and reward anticipation (Montoya et al. 2014), but also have detected timing-dependent changes in hippocampal memory functions. Specifically, when coinciding with declarative memory encoding, stress levels of CORT enhance long-term recall (Buchanan & Llova11, whereas their occurrence during retrieval impairs performance (de Quervain et al. 2000).

Most important, endogenous CORT and NE signals do not act in isolation, and there is accumulating experimental evidence that coactivation of both systems under emotional arousal is crucial for facilitating amygdala–hippocampus interplay during declarative memory formation. The resultant advantage of privileged declarative encoding of salient stimuli, however, comes at the expense of reduced recall of preceding and following information. This peri- emotional amnesia is BLA as well as β-noradrenergic dependent (Hurlemann 2006; Hurlemann et al. 2005; 2007a; 2007b; Strange et al. 2003) and further amplified, in both magnitude and temporal extent, by combined prelearning administration of exogenous CORT (30 mg po) and reboxetine (4 mg po), thus suggesting synergistic NE–CORT interactions (Hurlemann 2008; Hurlemann et al. 2007a). The same pharmacologic intervention was found to induce a negative response bias toward fearful faces in the centromedial nucleus of the amygdala (CMA), an effect that was absent when CORT levels were augmented alone (Kukolja et al. 2008). Evidence indicates that response shifts mediated by CORT, NE, and their interactions are not restricted to the CMA, but propagate to interconnected areas including the dorsal striatum, which can be prevented by blockade of mineralocorticoid receptors with spironolactone (400 mg po) (Vogel et al. 2015b).

Collectively, these findings argue for a reallocation of neural resources as a function of CORT and NE coactivation under emotional arousal, hence enabling prioritized access to the salience network and memory stores. Obviously, this mechanism confers costs and benefits, evident in a larger deviation of amygdala–hippocampal resources during encoding (Kukolja et al. 2011) and deactivation of prefrontal cortex (PFC) (van Stegeren et al. 2010). It has been conceptualized that such co-occurrence of deficient top-down control from PFC and enhanced amygdala–hippocampus interactions under conditions of heightened CORT and NE release may result in hypermnesia for emotional events, which, when manifest in extreme forms, is pathognomonic of post-traumatic stress disorder (PTSD) (Hurlemann 2008). Converging support for this etiologic model comes from preclinical (Bryant et al. 2013) and clinical studies (Nicholson et al. 2014), both of which suggest that CORT and CORT co-activation predisposes to the development of indelible memories. Future research addressing the mechanistic underpinnings of arousal-induced memory distortions in PTSD should, therefore, not only focus on neurotransmitter interactions between GLU and NE, as outlined by the GANE model, but also take the interplay of NE and endocrine players including CORT into perspective, which promotes stress-induced remodeling of neural architecture through (epi)genetic modifications as well as rapid non-genomic adaptations (de Kloet et al. 2005; McEwen et al. 2015). The latter include the non-genomic modulation of hippocampal glutamate transmission via activation of mineralocorticoid receptors (Karst et al. 2005), further illustrating the rapid susceptibility of memory functions to emotional arousal and stress.
Arousal-biased preferences for sensory input: An agent-centered and multisource perspective

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Abstract: I argue that the GANE (glutamate amplifies noradrenergic effects) model basically explains an arousal-based amplification of emotional stimuli, whereas effects on neutral stimuli indicate a contextualization process aiming to reduce stimulus ambiguity. To extend the model’s validity, I suggest distinguishing between internal and external emotional sources, as well considering the stimulus valence and addressing age-related differences in attention and memory preferences.

Mather et al. beautifully describe the neuronal mechanisms likely to account for an arousal-based modulation of selectivity phenomena in attention and memory. In addition to previous emotion-cognition models focusing on the competition between emotional and neutral stimuli, the GANE (glutamate amplifies noradrenergic effects) model also aims to explain how arousal resolves the competition between neutral stimuli. In this context, I propose a complementary perspective. I argue that the GANE model explains mainly the mechanisms underlying the processing of emotional stimuli, whereas co-occurring effects on the processing of neutral stimuli may be interpreted as side effects of a contextualization process targeting the emotional stimulus. The empirical evidence Mather and colleagues present suggests that arousal-based amplification or inhibition of neutral input is heavily constrained by the spatiotemporal relationship between an emotional stimulus, which is the driving source of arousal, and neutral stimuli constituting the sensory context. I claim that effects of emotional stimuli on proximal neutral ones are a signature of stimulus contextualization in favor of a disambiguation of emotional stimuli. Indeed, for the perceiving agent, emotional stimuli are often characterized by significant ambiguity (cf. Duval et al. 2013). For example, the recognition of an arousing facial expression may be context dependent (cf. Barrett et al. 2011). The valence and the discrete emotional category of an arousal stimulus are not completely inherent features of the stimulus, but they are also constituted by the context (including neutral stimuli) in which the arousing stimulus is embedded. Therefore, the processing of an emotional stimulus benefits from an amplification of salient neutral stimuli standing in an optimal spatiotemporal relationship to the emotional target stimulus. It appears beneficial for the human organism that emotional stimuli are not processed in isolation from rather neutral context information. Amplifying the processing of otherwise prioritized neutral stimuli in the presence of an emotional stimulus creates a context that facilitates the appropriate classification and encoding of the properties of the emotional stimulus. Indeed, the context seems to be routinely encoded during emotion perception (Barrett & Kensinger 2010). In this sense, the arousal-based mechanisms outlined in the GANE model cannot be generalized to settings in which emotional stimulation is rather negligible. However, based on this perspective, two aspects may help to further improve the conceptual framework of the model and its validity:

First, I propose that the GANE model would benefit from a more explicit distinction between internally and externally located sources of emotional arousal (cf. Kaspar 2013; Kaspar & König 2012). The current model addresses primarily the latter type, namely, sensory stimuli located outside of the perceiving agent, whereas internal forms of emotional arousal refer to the agent’s current emotional/mood state. While the arousing power of external stimuli is tied to the stimulus and, hence, places tight spatiotemporal constraints for an amplification or inhibition of the neutral surrounding, arousal elicited and maintained by internal thoughts of the agent might be more easily linked to any neutral stimulus. Of course, an internally located source of emotional arousal might be elicited by an external source, but some residual arousal (i.e., mood) continues for a while after source offset. Indeed, excitation-transfer theories (Bryant & Miron 2003; Zillmann 1983) propose that the residual arousal from a stimulus can be transferred to a subsequent stimulus, whereas the emotional valence of the stimuli may differ. If residual arousal can actually be tapped by a stimulus other than the original in this way (still to be shown), we can extend the validity of the GANE model by implementing a multisource approach to arousal-biased information processing.

Second, the model neglects the valence aspect of arousing sources being of central relevance, not only from the perspective of disambiguation tendencies. It seems that the arousal and valence ascribed to a stimulus are not completely independent features (cf. Kaspar & König 2012), whereby negative (vs. positive) stimuli show a tendency toward higher arousal (Ito et al. 1998; Kim & Hamann 2007). Thus, negative stimuli may have a higher potential to ignite neuronal hotspots and to fine-tune priority signals. This bias is plausible from an evolutionary perspective, as it is more prejudicial to miss a potential threat than to miss a potential reward. Thus, across different scenarios, external negative (vs. positive) stimuli may elicit stronger modulation effects on neutral stimuli being in an optimal spatiotemporal distance, whereas a negative (vs. positive) mood state might have more long-lasting effects because of more residual excitation.

Indeed, the location (internal vs. external) and valence of the arousing source have a critical influence on attention and memory processes. For example, younger adults showed an attentional preference and better memory performance for negative stimuli compared with positive stimuli, and this bias was more pronounced when participants had been in a positive (vs. negative) mood (Kaspar et al. 2015). Thus, internal and external sources of emotional arousal may show specific interactions depending on their valence. With respect to the GANE model, negative stimuli presumably have a higher likelihood to bias perception and memory when they are in strong contrast to emotional background noise within the perceiving agent. However, preferences for positive over negative information have also been reported for younger adults in specific circumstances (Becker & Leinenger 2011; Parrott & Sabini 1990; Schwager & Rothermund 2013), indicating more complex mechanisms of the processing of emotion-laden stimuli than delineated in an exclusive arousal-based model.

Finally, in this context, the GANE model suggests a brain-based explanation for the very reliable information processing bias toward positive (vs. negative) information in older adults (Reed et al. 2014). Mather and colleagues proposed that arousal may not increase selectivity similarly effectively among older adults because of age-related changes in the locus coeruleus–norepinephrine system. If so, negatively valenced stimuli may gradually lose their high arousing potential across the life span, facilitating controlled attentional shifts toward positive stimuli at an older age (cf. Hahn et al. 2006; Rought et al. 2007). Thus, the GANE model adds a brain-based explanation for this age-dependent change in biased competition that is discussed mainly in terms of the socioemotional selectivity theory (Carstensen et al. 2003) emphasizing age-related changes in emotion-regulation motivation.

Does arousal enhance apical amplification and disamplification?

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Abstract: We summarize evidence that input to the apical tufts of neocortical pyramidal cells modulates their response to basal input. Because this apical amplification and disamplification provide intracortical mechanisms for prioritization, Mather and colleagues argue that their effects are enhanced by noradrenergic arousal. Though that is likely, it has not yet been adequately studied. Their article shows that it should be.

Mather et al. argue that as arousal increases, things of high priority are perceived and remembered even better, whereas things of low priority are suppressed even more. Intracortical mechanisms for prioritization of selected signals are a prerequisite for this because the noradrenergic system provides only diffuse low-bandwidth innervation of neocortex, whereas the particular signals to be amplified or suppressed must be specified by locally specific interactions of high bandwidth. We therefore outline recent evidence for intracellular and microcircuit mechanisms by which signals are either amplified or suppressed within neocortex, prior to their further modulation by the noradrenergic system. We refer to these mechanisms as apical amplification (AA) and disamplification.

Evidence for AA is provided by patch-clamping studies showing that inputs to the apical tufts of pyramidal cells are integrated separately from inputs to their basal dendrites before being used to modulate the cell’s response. Current models of neocortex, including noradrenergic effects, typically assume that pyramidal cells can be adequately thought of as point processors that simply sum all of their excitatory and inhibitory inputs and fire when that sum exceeds a threshold. The evidence for AA indicates that some pyramidal cells have not one, but two main sites of integration such that when apical and basal inputs coincide, intracellular calcium spikes initiated by a site of integration near the top of the apical dendrite amplify the cell’s response to its basal inputs (Larkum 2013; Larkum et al. 1999; 2007; 2009).

The most studied mechanism by which AA is implemented in layer 5 is referred to as backpropagation-activated calcium spike firing (BAC firing). In addition to these two main integration sites, local integration takes place within both basal and tuft dendrites by the regenerative activation of N-methyl-D-aspartate receptors (NMDA spikes). AA may be fully implemented by NMDA spikes alone in supragranular neurons (Palmer et al. 2014), but even in subgranular neurons, NMDA spikes have an important influence (Larkum et al. 2009). Essential properties of these mechanisms are illustrated in Figure 1. Inhibitory interneurons that specifically target apical dendrites in layer 1, such as Martinotti cells, produce disamplification, which suppresses amplification without inhibiting action potential output.

Though much of this work has been carried out in vitro, there are strong grounds for supposing that AA and disamplification apply to awake behaving animals (Phillips et al. 2013). Imaging studies of local dendritic NMDA spikes in awake behaving animals indicate the importance of such integrative intracellular processes in vivo (Cichon & Gan 2015; Gambino et al. 2014; Grienberger et al. 2014; Lavzin et al. 2012; Palmer et al. 2014; Smith et al. 2013; Xu et al. 2012). These discoveries are well known to cellular neurophysiologists, but not yet to psychologists or cognitive neuroscientists. For a clear introduction to AA and disamplification and their relevance to cognitive function and theoretical neuroscience, see Phillips (2015).

Arousal releases norepinephrine (NE), that is, noradrenaline, which regulates the firing mode of layer 5 neurons (Wang & McCormick 1993). Many new questions are raised by the possibility of interactions between AA and NE release in these and other neocortical neurons. First, are the effects of NE and AA synergistic, or do they simply sum in some quasi-linear way? Synergistic interactions between AA and mechanisms proposed in the GANE model seem likely because glutamate spillover will not spread from apical to basal dendrites. Spillover is intrinsic to the GANE model because of the non-synaptic component of NE release, and that implicates NMDA more than AMPA receptors. Local dendritic NMDA spikes are also enhanced by glutamate spillover (Chalfoux & Carter 2011). To see the possibility of synergistic interactions consider the case to which AA is most applicable, that is, where apical input is strong and basal input is present but weak. There would then be NE-dependent enhancement of depolarization in the apical tuft but not in the basal dendrites. That would increase the effect of AA on cellular output while maintaining the need for basal input to initiate axonal spiking. Second, how are NE-receptor subtypes distributed across regions, layers and subcellular compartments, and is that compatible with the modulatory role proposed for tuft inputs? An explicit focus on intracellular and microcircuit mechanisms in theories of arousal requires answers to these questions.

![Figure 1](https://www.cambridge.org/core/figure/18b6c66fdd8c0bdf0f722c482f36e907/fig1)[Figure 1](https://www.cambridge.org/core/figure/18b6c66fdd8c0bdf0f722c482f36e907/fig1) (Larkum & Phillips). Dendritic spikes in neocortical pyramidal neurons. Apical tufts of pyramidal neurons receive inputs from diverse sources. Calcium currents, and thus synaptic plasticity, depend on backpropagating action potentials (bAPs, gray), apical dendritic calcium spikes (red) and NMDA spikes (blue). NMDA spikes require both local depolarization and glutamate (blue dots) and are enhanced by glutamate spillover to extrasynaptic NMDA receptors (green squares). Norepinephrine (maroon dots) interacts with glutamate in a feedback process hypothesized to enhance post-synaptic excitability.
Third, will studies of interactions between AA and NE cast light on the putative role of AA in regulating states or levels of consciousness (Bachmann & Hudec 2014; Meyer 2015; Phillips 2015)? It seems likely that they will. Fourth, do previous studies under-estimate the extent of AA because they do not ensure appropriate levels of noradrenergic input? This is clearly relevant to in vitro studies or under anesthesia, but, Mather and colleagues’s hypotheses imply that local phasic arousal needs to be considered as well as tonic arousal when studying awake behaving animals. Finally, are working memory capabilities dependent upon specialized interactions between NE and AA in dorsolateral prefrontal cortex (Arnsten et al. 2012)?

Much intracellular, electrophysiological, cognitive, and computational research is required to answer such questions. If that shows noradrenergic enhancement of AA and dissimilification, then that will strengthen and broaden both the GANE model and our understanding of the role of intracellular computations in mental life. If not, then we will need to discover why not. Thus, the target article opens the door to a wide array of issues concerning interactions between noradrenergic arousal and prioritization within the neocortex by AA and dissimilification. These issues may well be crucial to our understanding of relations between brain and behavior.

**Emotional memory: From affective relevance to arousal**

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**Abstract:** Arousal is typically conceived as a key component of emotional response. We describe here the psychological processes thought to elicit arousal—in particular, the processes involved in the appraisal of affective relevance. The key role of relevance in attentional and memory processing, and its links with arousal, is discussed with respect to the GANE (glutamate amplifies noradrenergic effects) model described by Mather et al.

Mather et al. provide an innovative and integrative model aimed at explaining, at the neural level, how arousal can both enhance and impair cognitive processing, such as perception and memory. The glutamate amplifies noradrenergic effects (GANE) model proposed by the authors accounts for results indicating that increased norepinephrine under arousal effects prioritization of information processing, for example, enhances memory for salient information at the expense of mundane information. Mather et al. consider arousal to be the critical factor that amplifies the perception of emotional stimuli while impairing the perception of other concurrent stimuli, as has been proposed in the arousal-biased competition (ABC) theory (Mather & Sutherland 2011). Mather et al. are particularly interested in analyzing the effect of selectivity in the perception and memory of emotional stimuli under arousal; however, the construct “arousal” is not conceptually clear if one considers how it is used in the affective sciences literature, especially concerning theories of emotion. More generally, the relationship between emotion and arousal is far from being consensual, notably when it comes to modeling the psychological mechanism that elicits different types of arousal (see Sander 2013). For example, Frijda (1986) distinguished among three response systems that embrace the construct of arousal or activation: autonomic arousal, electrocortical arousal, and behavioral activation. Traditionally, theories of emotion refer to arousal mainly in terms of either (1) the consciously felt activation dimension of the experienced affect or (2) the bodily reaction during an emotional episode, specifically the sympathetic nervous system. Dimensional theories propose that any experienced affect can be analyzed as a bodily feeling on a dimension of an activated (e.g., as in surprise) or deactivated (e.g., as in boredom) state (Feldman 1995; Russell 1989). The bodily reaction can be measured in terms of a psychophysiological state driven by the activation of the sympathetic nervous system. Emotion researchers typically use the construct of arousal to refer to the activation of the sympathetic nervous system, and physiological measures (e.g., skin conductance response or pupil dilation) are often used as measures of emotion intensity, as they are also considered to mirror subjective affective ratings (see Bradley et al. 2008; Lang et al. 1993).

The effect of arousal on performance has been revealed in various research domains and is consistent with the model proposed by Mather et al., suggesting that—under arousal—neutral stimuli with a high priority are processed preferentially.

The priority factor is essential to memory enhancement. Indeed, some have proposed that arousal per se is not sufficient to explain memory improvement for neutral information (Reisberg & Heuer 2004). For example, a study that manipulated arousal by injection of adrenalin versus saline showed no difference in the results of memory performance for verbal descriptors (e.g., name, occupation) presented previously with neutral faces, although heart rate and skin conductance increased in the group with adrenalin injections (Christianson & Mjörndal 1985). The information presented did not differ in terms of priority. This notion of priority, especially when related to the well-being or major concerns of the individual, is considered to be key in explaining emotion elicitation. Indeed, appraisal theories of emotion suggest that stimuli that are appraised as relevant for an individual’s concerns (e.g., that relate to goals, needs, and values) are typically those that have the competency to elicit an emotional response by driving changes in action tendencies, in expressions, in the peripheral nervous system, and in the conscious feeling (e.g., Sander et al. 2005). With respect to arousal, a conceptually interesting advantage of such models is that an explanation for the elicitation of arousal is proposed: only concern-relevant events elicit arousal. Evidence indicates that stimuli that are appraised as concern relevant not only elicit such emotional responses, but also are prioritized in attention (see Pool et al. 2015). For example, a recent meta-analysis revealed that the magnitude of the attentional bias for positive stimuli varies as a function of arousal, but also that this bias is significantly larger when the stimulus is relevant to specific concerns (e.g., hunger) of the participants compared with other positive stimuli that are less relevant to the participants’ concerns (Pool et al. 2015). Interestingly, in this meta-analysis, when arousal and relevance moderators were tested by statistically controlling their respective variances, only relevance remained a significant predictor of the magnitude of this bias in emotional attention. Emotional attention is a process that has been strongly rely on the amygdala (Vuilleumier 2005a), a region that has also been suggested to be critical for the process of appraised relevance (e.g., Cunningham & Brosch 2012; Sander et al. 2003) and of importance in promoting memory consolidation, as described by Mather et al. (see Roozendaal & McGaugh 2011 for a review).

Studies supporting the ABC theory (e.g., Sutherland & Mather 2012) and GANE models are based largely on experimental manipulation showing that high-priority neutral information under external arousal is enhanced in memory (as compared with low-priority neutral information). Showing that goal-relevant events, which are intrinsically neutral, are processed preferentially might extend the model proposed by Mather et al. For example, the goal relevance hypothesis of memory facilitation has been

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tested in a gamelike study in which the goal of the participant was to win points. Participants showed better memory performance for initially neutral items that signaled a gain (i.e., goal-conducive items) than for initially neutral items that were goal irrelevant (Montagrin et al. 2013).

Our suggestion is that events that are relevant to one’s goals elicit an arousal response, capture attention, and facilitate memory. A fascinating research topic, for which both the GANE model and the ABC theory can be particularly well articulated with appraisal theories of emotion, is to understand the causal relationships between relevance detection, arousal, attention, and memory.

Effect of arousal on perception as studied through the lens of the motor correlates of sexual arousal

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Abstract: The study of sexual arousal is at the interface of affective and social neurosciences. Recent results regarding the motor correlates of sexual arousal demonstrating an early freezing response are in perfect accordance with the GANE (glutamate amplifies noradrenergic effects) model’s sustaining the double role of the arousal dimension on emotional processing.

During recent decades, sexual affiliation has been an exponentially explored functional context within socioaffective neuroscience, especially with respect to the motivational component of inter-attraction, which can be either positive or negative within social relationships. As appearing in international databases, sexual visual stimuli are often reported as the most arousing stimuli and are therefore ideal to increase knowledge of the influence of the arousal dimension on the neural (central and peripheral) and psychological correlates of emotional information processing, which is at the center of the theory presented by Mather et al. Here, we focus on recent results regarding the motor correlates of visual sexual information corroborating the complex modulatory role of arousal as developed in the GANE (glutamate amplifies noradrenergic effects) theory.

Within the framework of sexual behavior, emotion can be conceptualized partly as an action disposition characterized by a context-dependent (e.g., approach vs. avoidance) behavioral component, which may be mediated by automatic responses (Campbell et al. 1997; Panksepp & Biven 2012). In that sense, emotion should influence several steps of the motor response (Bradley et al. 1992; Hellbig et al. 2011; Williams et al. 1996) by inducing an approach behavior to what promotes our well-being and our survival and an avoidance behavior in response to painful experiences (Elliot & Covington 2001). Albeit recent, most neuroimaging studies of sexual arousal made central in their theoretical model a motivational component. Several

Figure 1 (Mouras). Means and standard deviations of postural indices as a function of the stimulus. (A) Amplitude of the sway of the center of pressure (COP) in the mediolateral direction (Amp [COP]-ML). (B) Amplitude of the sway of the COP in the anteroposterior direction (Amp [COP]-AP). (C) Standard displacement of the COP in the mediolateral direction (SD [COP]-ML). (D) Standard displacement of the COP in the anteroposterior direction (SD [COP]-AP). (E) Area encompassed by displacements of the COP (COP-area). *p < 0.05, **p < 0.01 when comparing stimuli. (From Mouras et al. 2015).
What do we GANE with age?

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Abstract: Mather and colleagues provide an impressive cross-level account of how arousal levels modulate behavior, and they support it with data ranging from receptor pharmacology to measures of cognitive function. Here we consider two related questions: (1) Why should the brain engage in different arousal levels? and (2) What are the predicted consequences of age-related changes in norepinephrine signaling for cognitive function?

Mather and colleagues have developed an impressive theoretical model linking arousal-mediated changes in cognition to the local signaling dynamics at the axon terminals of noradrenergic locus coeruleus (LC) projections. The authors provide compelling evidence for how this link between biology and cognition is made; however, they leave open a key question: Why should the brain undergo fluctuations in arousal that influence information processing? Answering this question is important not only for appreciating the intricate biological design proposed by Mather and colleagues, but also for understanding the contexts in which such a design would be maladaptive. Here we explore the why question and speculate that healthy aging may constitute one such maladaptive context.

So, if the brain has a good system for prioritizing sources of information, why should it ever be turned down? One possible answer is that prioritized sources of information are often imperfect predictors of behaviorally relevant variables. In such cases, inferring the variable of interest (e.g., the best foraging location) requires combining probabilistic sensory information (e.g., the color of the bush in a distant hill) with currently held beliefs (how many berries are expected based on past experience). The relative contributions of these factors should be determined by the precision with which they predict the behaviorally relevant variable; thus, sensory information should be discounted when prior expectations are strong (e.g., the forager has recently counted the berries on the bush) or when sensory evidence is weak (e.g., the color of the bush is a bad predictor of caloric yield). In either circumstance, amplifying the prioritized sensory information would disrupt inference by allowing poor sensory cues to overwhelm precise internal expectations. There is some evidence to suggest that arousal levels are decremented under such conditions. Pupil diameter, a marker for arousal, is large during periods of uncertainty and constrains as expectations become more reliable (Lavín et al. 2014; Nassar et al. 2012; Nieuwenhuis et al. 2011; Preuschoff 2011). Information that is inconsistent with expectations drives sharp increases in arousal that appear to affect the relative influence of new observations on behavior (Lavín et al. 2014; Nassar et al. 2012; Nieuwenhuis et al. 2011; Preuschoff 2011). These data suggest that decrements in arousal, likely implemented through reductions in the firing of LC neurons, may serve an important role in optimal inference (Joshi et al. 2016). In particular, one role of low arousal levels might be to protect strong internal predictions from prioritized but potentially distracting information.

Another normative justification for reducing the influence of priority maps is that under some conditions, it is useful to explore alternatives to the current course of action that might provide better long-run returns. For example, information about a known source of reward (e.g., the berry bush on the hill) might be prioritized over other potential sources of reward that are yet to be discovered (e.g., an apple tree that only recently began to bear fruit). Thus, reducing the influence of priority maps could provide an incentive to explore non-prioritized inputs. Exploring potential alternative reward streams becomes important as the known source of reward is depleted, particularly if there is sufficient time to capitalize on any knowledge gained in the exploratory process (Wilson et al. 2014). There is some evidence to suggest that shifts from exploiting a known source of reward to exploring alternative options are accompanied by a shift from a phasic (stimulus evoked) mode of LC firing to a high tonic mode (Aston-Jones & Cohen 2005; Jepma & Nieuwenhuis 2011). Thus, fluctuations in arousal might allow for an effective navigation of the trade-off between exploitation and exploration in addition to optimizing inference.

The biological mechanism proposed by Mather and colleagues suggests that the optimization of inference and exploration
through fluctuations in arousal may be highly sensitive to the state of the LC–norepinephrine (NE) system and its biophysical components. There is some evidence for dysregulation of this system over the course of healthy aging. Findings from histologic post-mortem studies point to a substantial cell loss in the LC with age (Chan-Palay & Asan 1980; Grudzien et al. 2007; Manaye et al. 1995). Moreover, neuronal density in LC is strongly related to cognitive decline in the period before death (Wilson et al. 2013). These findings seem to line up with recent in vivo structural magnetic resonance imaging findings that point to neuromelanin-related magnetic resonance signal loss with age (Shibata et al. 2006). Taken together, these findings suggest that aging is associated with substantial structural changes in the LC, which are associated with cognitive decline.

One potential cause of age-related cognitive decline could be that this pattern of changes in the NE system disrupts optimal inference. In particular, lower levels of NE could prevent the positive feedback of glutamate on NE release from achieving high enough NE levels to activate low-affinity β-adrenoceptors proposed by Mather and colleagues. This could lead to a suppression of high-priority signals, even at high arousal levels associated with uncertainty, when new information should be highly influential on behaviorally relevant beliefs. Consistent with this notion, older adults show selective behavioral impairments at learning under conditions of uncertainty, the same conditions that typically drive increased arousal and increased influence of new information on learning in younger participants (Eppinger et al. 2008; 2011; Nassar et al. 2012; 2016). The cause of this learning impairment has not been directly linked to the function of the LC–NE system to date, but in light of the biological link provided by Mather and colleagues, it should be explored in the very near future.

Changes in NE functioning with age may also affect the ability to mediate exploration and exploitation in older adults. In particular, reduced NE levels may prevent phasic signals from activating β-adrenoceptors, even in regions where signals are prioritized through association with an explorative action. This would reduce the contrast between exploitative and exploratory action representations and shift behavior toward a more exploratory regime. Interestingly, increased choice variability, which can be used as a strategy for random exploration, is enhanced in older adults across a wide range of tasks (Garrett et al. 2013). Future work should focus on animal models where the mechanisms for age-related changes can be explored at the level of detail specified by Mather and colleagues.

Mather et al. have proposed an intriguing theory to explain how norepinephrine (NE) release and subsequent noradrenergic modulatory actions are focused in neural circuits by concomitant “priority” stimulus-driven release of glutamate. In doing so they confront a question that has perplexed the field for some time, that is, how to account for selectivity of signal processing in noradrenergic terminal fields and focused perception of salient events while maintaining broadband discharge signaling. This nucleus locus coeruleus (LC) is elevated, as would occur during generalized arousal. Here we focus on how the theory applies to NE modulation of sensory signal processing. Given the results of four decades of published work, we would expect increases in LC–NE output to promote enhanced neuronal and neural network responses to sensory-driven afferent inputs (Berridge & Waterhouse 2003), actions that have been linked to improved performance of sensory-guided behavioral tasks (Aston-Jones et al. 1999; Rajkowski et al. 2004).

Until recently, conventional wisdom was that the LC–NE efferent network was broadly distributed from a relatively small number of brainstem neurons with homogeneous physiological properties. Given this, the presumption was that NE was released uniformly and simultaneously across all terminal fields within the forebrain, cerebellum, and spinal cord for as long as LC is discharging, either tonically or phasically. If that were the case, neuronal and neural circuit responsiveness to sensory driven afferent inputs would be increased throughout the central nervous system without any bias in favor of one sensory signal versus another. If responsiveness to synthetically driven inputs is elevated everywhere and for every modality, what has been gained? Is there a way for the LC–NE system to selectively differentiate sensory signals from the constant stream of information that is presented to the nervous system from the periphery? Mather and colleagues’ GANE (glutamate amplifies noradrenergic effects) theory is timely in so far as it appropriately confronts these issues.

An idea similar to the current theory was suggested by Marrocco et al. (1987) after they observed a correlation between catecholamine release in monkey visual cortex and coincident light-evoked activity in geniculostriate projections to ocular dominance columns. These authors postulated a local interaction between NE fibers and geniculostriate afferents: one that created a local hotspot for NE release within the visual cortex and, thus, preferentially promoted modulation of synaptic transmission at this site. Akin to Marrocco and colleagues’ proposal, the GANE theory argues that locally released glutamate provides the means for amplifying release of NE from tonically or phasically active LC–NE fibers.

The GANE theory accounts for many, but not all of the well-documented attributes and operational capacities of the LC–NE system, particularly those demonstrated in sensory networks. The authors exhaustively reviewed an extensive literature including many reports that support the core of their proposal: a positive feedback mechanism whereby synaptic release of glutamate amplifies NE release from nearby noradrenergic axons and results in enhanced responsiveness of neurons and glia to glutamate neurotransmission at this local site of interaction. The process relies on a delicate balance and interplay between receptor-mediated actions that are triggered as extracellular concentrations of NE and glutamate change. The temporal and spatial dynamics of these interactions are postulated, but experimental evidence to support the details of these mechanisms is lacking. For example, to date, the extracellular tissue concentrations of NE that yield the range of modulatory actions demonstrated in vivo and in vitro have been only crudely approximated based upon the results of microdialysis studies (Berridge and Abercrombie 1999; Florin-Lechner et al. 1996). As illustrated in many studies, LC–NE modulatory effects are expressed according to an inverted-U dose–response curve, rising to optimal facilitation of cellular and behavioral events as LC–NE activity increases and then falling to suppression of neuronal responsiveness and disrupted task performance as NE concentrations and LC
Commentary/Mather et al.: Norepinephrine ignites local hotspots of neuronal excitation

Competition elicits arousal and affect

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Abstract: The emotion–cognition integration in Mather et al. can be extended by specifying the relationship between competition and arousal in the reverse direction. According to affective monitoring, competition raises arousal, which, when sustained, results in negative affect, evoking theta oscillations, and when resolved, in positive affect, evoking gamma oscillations. Competition should be considered a core process in both cognition and emotion.

Competitive processes in the brain have the potential to account for a much larger range of behavioral functions than only attention and memory. Mather et al. successfully account for enhancing and impairing effects of arousal (and norepinephrine [NE]) on selective attention and memory in terms of neural competition, but still sometimes mix the brain metaphor with the “steam engine” metaphor. If competition is envisaged as mutual inhibition between neural nodes, there is no further need to borrow conservation laws from nineteenth-century physics and invoke limited resource models. When arousal effects are thus predominantly analyzed in terms of neural processes, the source of arousal and emotion is discussed here only on a behavioral level. The analysis can be extended by also grounding arousal and affect in processes of neural competition.

Emotion research abounds in examples where modulatory influences of affect on attention and memory appear to be reciprocal, in that a similar, but not affectively laden, manipulation of attention or memory is able to elicit affect (e.g., Dreisbach & Fischer 2012; Phaf & Rotteveel 2005; Rotteveel & Phaf 2007; Srinivasan & Hanif 2010). Most relevant to the present discussion may be studies demonstrating distracter devaluation, and target appreciation, in attentional filtering tasks (Goolsby et al. 2009; see also Raymond et al. 2003). The present authors explain target selection in attentional tasks by biased competition (Desimone & Duncan 1995; Duncan 1996; see also Phaf et al. 1990), so the appreciation and devaluation may also follow from competitive processes. Phaf and Rotteveel (2012) have argued that affect is the consequence of competition (i.e., conflicting neural representations [cf. Murre et al. 1992]), and that negative affect arises when it is sustained, but positive affect when it is quickly resolved. Fluent, competition-less processing in itself is not sufficient to raise positive affect (but see Fazendeiro et al. 2007). The common denominator in both positive and negative affect is the initial competition, which may thus correspond to arousal. The largely similar effects of positive and negative arousal (e.g., Sutherland & Mather, under review) may thus result from this initial phase of competition.

Gamma and theta oscillations are involved but not well integrated by the authors in their competitive framework. Such integration, however, serendipitously emerged from our evolutionary simulations (Heerebout & Phaf 2010a, 2010b). Random variation combined with selection of the fittest individuals led to the development of both competition and oscillations in neural networks that controlled agents roaming an artificial environment. The fitness measure combined the amount of food gathered and the time the agent managed to escape from predators. The serendipitous emergence of oscillations coincided with a near doubling of fitness, indicating that they were very functional to the agents. In fact, the same feedback loops between excitatory and inhibitory nodes developed autonomously in the evolutionary simulations as were suggested in Mather et al. Heerebout and Phaf (2010a) investigated the behavior of these agents and found that the function of oscillations was complementary to that of competition. Competition enables the selection of representations, and oscillations allow for a reset of winners (i.e., switching of representations).

Attentional flexibility is more functional with positive than with negative affect. When searching for food, it is highly adaptive to be able to quickly shift attention toward an approaching predator. When running from a predator, it is highly maladaptive to switch attention to some palatable food (i.e., “it is better to miss dinner than to be dinner”). We hypothesized that low-frequency, presumably theta, oscillations are evoked by competition, whereas high-frequency, presumably gamma, oscillations arise when the competition is solved. Similar to arousal (and NE), the former increase selectivity and narrow attentional focus, whereas the latter enhance switching between representations and attentional flexibility (Bauer et al. 2009; Heerebout et al. 2013). Crucially, in this view, theta oscillations (i.e., resulting from competition) precede gamma oscillations and may thus be the primary reflection of arousal. According to this framework, without competition there is no arousal, theta synchrony (i.e., negative affect), or eventual gamma synchrony (i.e., positive affect).
Adaptive memory systems for remembering the salient and the seemingly mundane

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Abstract: In an adaptive memory system, events should be prioritized in memory based on their own significance, as well as the significance of preceding or following events. Here we argue that tag-and-capture models complement the GANE (glutamate amplifies noradrenergic effects) model by describing a mechanism that supports the transfer of memory benefits from one event to the next.

Imagine you are enjoying a brisk hike through the forest. You round a bend and stop dead in your tracks—a large bear is on the trail ahead, staring directly at you. Your attention is entirely focused on this unexpected and potential threat. You remain unharmed, but you will remember this moment for years to come. The GANE (glutamate amplifies noradrenergic effects) model provides a compelling account of how arousal at the moment of experience leads to selective memory for prioritized information—be it the bear, or the experience of emotional events, however, we often remember other details that seemed unimportant at first, but were experienced in connection to the emotional event. For example, you might also remember seeing a fresh animal print in the mud earlier in your hike. These memories are adaptive; you do not want to wander unprepared into bear territory again. How we selectively remember information that occurred minutes to hours before an emotional experience is outside of the scope of the GANE model, but is well explained by a tag-and-capture model of memory consolidation.

Tag-and-capture refers to a model by which memory traces that are tagged during learning can benefit from periods of enhanced plasticity prior to or after learning, by capturing the plasticity-related products (PRPs) necessary for long-term consolidation (Redondo & Morris 2011; Viola et al. 2014). A key feature of this model is that weakly encoded memories stand the most to gain from this form of modulation, in that they are insufficient to drive long-term consolidation on their own. Moreover, the tag and capture phases need not occur simultaneously but can be separated by minutes to hours as long as they affect the same neural targets. Although tag-and-capture models were initially applied to electrophysiological studies of long-term potentiation (Frey & Morris 1997; 1998), it has since been shown that salient or arousing experiences, such as novelty exposure, can rescue weak memories (Moncada & Viola 2007; Wang et al. 2010) that overlap with the salient event (Ballarini et al. 2009).

A critical distinction between the GANE and tag-and-capture models is the time scales on which they are expected to operate. The GANE model proposes simultaneous engagement of noradrenaline and glutamate systems to enhance memory. Because this model necessitates coincidence detection across these neurotransmitter systems, the time frame by which arousal can facilitate learning of non-prioritized information is limited to the duration of salient memory’s overlap with a separate arousing event. It is worth noting that both sets of mechanisms can, in theory, be deployed at any site of plasticity, offering flexibility in terms of which learning systems can benefit from arousal.

The relative temporal flexibility of tag-and-capture results from mechanisms that are distinct from GANE, including dopaminergic neuromodulation (Redondo & Morris 2011). Critically, the dopaminergic system has properties that allow it to support consolidation at extended time scales. First, dopamine release in response to arousal is characterized by tonic, as opposed to phasic, activation (Grace et al. 2007), such that a single arousing event could result in prolonged increases in dopaminergic tone and facilitated learning (Shohamany & Adcock 2010). Second, dopamine acts on relatively late stages of memory consolidation, allowing for salient events and encoding to be disparate in time. That is, dopamine affects protein synthesis-dependent long-term potentiation—a process necessary for consolidation—rather than to oppose memory encoding via early long-term potentiation (Lisman et al. 2011). Because dopamine-mediated synthesis of PRPs can occur independently from encoding, it may be particularly relevant for the consolidation of weakly encoded events, relative to strongly encoded events that are able to initiate PRP synthesis on their own through mechanisms...
Importance of amygdala noradrenergic activity and large-scale neural networks in regulating emotional arousal effects on perception and memory

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Abstract: Mather and colleagues postulate that norepinephrine promotes selective processing of emotionally salient information through local “hotspots” where norepinephrine release interacts with glutamatergic activity. However, findings in rodents and humans indicate that norepinephrine is ineffective in modulating mnemonic processes in the absence of a functional amygdala. We therefore argue that emphasis should shift toward modulatory effects of amygdala-driven changes at the network level.

Emotional arousal enhances memory of currently relevant—that is, salient—information, whereas it can impair memory of irrelevant information (Berman et al. 2013; Mather & Sunderland 2011). Mather et al. formulate the interesting hypothesis that when norepinephrine (NE) release coincides with high glutamatergic activity within an activated brain region or neuronal ensemble, NE release is increased further, resulting in locally enhanced neuronal activity and better memory. In contrast, when NE release does not coincide with high glutamate levels, NE suppresses neuronal activity, resulting in memory impairment. Although their model incorporates interactions at the systems level, it places strong emphasis on local processes, creating NE “hotspots.” Here, we argue that such primarily local effects underestimate the importance of modulatory influences of the amygdala on encoding and consolidation of information throughout the network and that, without a functioning amygdala, such NE hotspots might be unable to affect local mnemonic processes.

According to the widely accepted “amygdala modulation hypothesis,” basolateral amygdala (BLA) activity enhances memory of emotionally arousing experiences by influencing neural plasticity mechanisms in target regions elsewhere (McGaugh 2002). In rodents, pharmacologically enhancing or reducing noradrenergic activity within the BLA, that is, mimicking different arousal conditions, is sufficient to alter training-associated neural plasticity in distal brain regions (Beldjoud et al. 2015; McIntyre et al. 2005) and to determine whether neural representations in these regions are being strengthened (Roozendaal & McGaugh 2011). Recent evidence suggests that such BLA interactions with other brain regions not only modulate the strength of memory, but also are significantly involved in regulating memory precision (Ghosh & Chattarji 2015), and that NE activity in particular may be the driving force behind improved accuracy (Barsegyan et al. 2014). Human neuroimaging research corroborates these findings by showing that amygdala activity during encoding of emotionally arousing stimuli predicts enhancement of hippocampus-dependent memory (Canli et al. 2000; Hamann et al. 1998). β-Adrenergocortical blockade during encoding abolishes the emotional memory enhancement effect (Cahill et al. 1994) and suppresses memory-related amygdala activity (Strange & Dolan 2004). Amygdala–hippocampal connectivity, furthermore, is stronger for emotionally arousing than for neutral stimuli (Dolcos et al. 2004), and the dominant directionality of this connectivity is indeed from amygdala toward hippocampus (Fastenrath et al. 2014).

Critically, amygdala–NE interactions selectively enhance memory for emotionally arousing as compared with neutral stimuli (e.g., Cahill et al. 1994). Mather et al. posit that the amygdala modulation hypothesis explains this selectivity in terms of a trade-off in which resources are shifted toward the emotional stimuli. However, recent findings indicate that there may be more to it than a simple trade-off. For example, Lovitz and Thompson (2015) report that intra-BLA infusion of the β-adrenoceptor agonist clenbuterol induces a long-term increase in excitability of hippocampal neurons when administered after emotionally arousing inhibitory avoidance training, but that clenbuterol decreases hippocampal excitability in non-trained control animals. These findings strongly support the idea that the impairing effects of amygdala–NE interactions on memory of non-salient/non-arousing information involve an active process that is dependent on the amygdala.

Converging human evidence for this notion comes from patients with damage to the amygdala. For example, patients with Urbach–Wiethe disease (UWD), who exhibit selective calcifications in the BLA (Terburg et al. 2012), fail to show emotional enhancement of episodic memory (Cahill et al. 1995). Furthermore, studies in patients with other forms of amygdala pathology revealed a deficit in upregulating processing of emotional stimuli in higher-order visual cortices (Vuilleumier et al. 2004), as well as an impairment in increasing encoding-related hippocampal activity for emotional items (Richardson et al. 2004). Critically, UWD patients also exhibit enhanced memory for neutral information encountered in close temporal proximity to emotionally arousing stimuli (i.e., diminishing the impairment for such information observed in healthy controls [Strange et al. 2003]). One could argue that such findings remain consistent with an interpretation in terms of local hotspots of NE activity if amygdala damage would lead to a general impairment of NE signaling. However, UWD patients, although they fail to acquire conditioned responses, appear to exhibit normal arousal responses, as evidenced by normal skin conductance and startle responses to unconditioned stimuli (Bechara et al. 1995; Klumpers et al. 2015). Thus, findings from amygdala-lesioned patients agree with work in animals in suggesting that because of BLA damage, NE is ineffective in modulating local memory processes elsewhere in the brain.

Other studies have indicated that stress-related hormones such as glucocorticoids also contribute to selective enhancement of mnemonic memories. For example, in humans, elevating stress hormone levels after learning generally leads to consolidation benefits for emotionally arousing as compared with neutral information.
Locus coeruless reports changes in environmental contingencies

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Abstract: The GANE (glutamate amplifies noradrenergic effects) model proposed by Mather et al. attempts to explain how norepinephrine enhances processing in highly activated brain regions. Careful perusal of the sparse data available from recording studies in animals reveals that noradrenergic neurons are excited mainly by any change in the environment—a salient, novel, or unexpected sensory stimulus or a change in behavioral contingencies. This begets the “network reset hypothesis” supporting the notion that norepinephrine promotes rapid cognitive and behavioral adaption.

The functional significance of neuronal activity in a particular brain region or population of neurons is found in the environmental stimuli or cognitive context that drive (or inhibit) activity in those neurons. Thus, we know from electrophysiological exploration that the function of primary visual cortex is to respond to light, that of auditory cortex, to sound, and so forth. Thalamic nuclei have likewise been delineated in terms of function. The role of prefrontal cortex in working memory was hypothesized by lesion studies, but clearly demonstrated by recording neuronal activity in monkeys performing working memory tasks (Fuster 1991; Goldman-Rakic 1990). Likewise, single unit recording in rats performing spatial navigation tasks established the fundamental role of the hippocampus in spatial cognition (O’Keefe & Dostrovsky 1971). One of the principal functions of neurons of the ventral tegmental area is to report reward prediction error, based on recordings from this region in monkeys performing operant tasks. Thus, to gain a full understanding of the functional role of the locus coeruless–noradrenergic system (LC–NE), it is important to carefully consider what drives this small population of neurons. Until recent biotechnological developments, the only way to achieve this was through recording activity of LC neurons in unanesthetized animals in carefully controlled behavioral situations. Given the inaccessible pontine location and very small size of this nucleus, the task has proved to be challenging and the resultant literature quite sparse. Nevertheless, there are some studies that provide insight that goes beyond LC–NE mediation of arousal and response to salient, stressful, or novel stimuli, as summarized by Mather et al. in section 4.1. Furthermore, recent advances in functional magnetic resonance imaging resolution have allowed imaging of this nucleus in humans performing complex cognitive tasks. These studies are now corroborating a role for LC in cognitive flexibility and behavioral adaptation, although neurophysiological studies in animals and humans are still sparse.

The earliest recordings of activity of LC in behaving rats established its role in vigilance and its responses to salient environmental stimuli in all modalities (Aston-Jones & Bloom 1981). Subsequent experiments in rats and monkeys showed that LC neurons display remarkable plasticity as a function of environmental contingencies. Sensory responses habituate after just a few repetitions, even when initially robust, when no behavioral adaptation is required (Hervé-Minvielle & Sara 1995). In a hole board environment, encounter with a novel object elicits a robust phasic burst of LC neurons that persists for only one or two subsequent investigations of the object (Vankov et al. 1995). Differential conditioning studies have shown that LC cells are exquisitely sensitive to stimulus–reward contingencies, showing task-related responses at the very earliest stages. At the beginning of training, both conditioned stimuli and primary reward elicit phasic responses in LC neurons. After just a few trials, response to reward disappears and response to the stimulus predicting reward (CS+) increases, whereas response to the neutral stimulus (CS−) decreases. These discriminative conditioned responses in LC appear many trials before any behavioral expression of learning and before task-related responses emerge in the prefrontal cortex. They are not maintained during overtraining, but when contingencies change abruptly, as during extinction or reversal training, phasic LC responses are immediately reinstated, tens of trials before behavioral expression of learning (Bouret & Sara 2004; Sara & Segal 1991). Similar phenomena have been reported for behaviorally contingent LC activity in monkeys (Aston-Jones et al. 1997; Rijksowski et al. 2004).

These relatively sparse data collected from behaving rats and monkeys over a span of 25 years led us to hypothesize that NE released in the cortex in response to a salient event or to a sudden change in environmental contingencies acts to facilitate or promote a rapid change in cortical state, “reset” the active network, and drive an adaptive behavioral response (Bouret & Sara 2005; Sara & Bouret 2012). We have provided some preliminary evidence for a “reset” action of NE, revealed by spike-triggered wave form averages of gamma filtered local field potential. Gamma band synchronization (GBS) has functional roles in diverse cognitive processes, including attention, stimulus processing decision making, and response timing (Bosman et al. 2014). We found a strong temporal relation between GBS and spontaneous LC bursts. In fact, LC spiking interrupts the gamma wave for about 200 ms, with the recovered GBS having increased power (Poe and Sara 2014; Sara 2015, Fig. 3).

Recent functional magnetic resonance imaging studies in humans have lent strong support to a prediction of Corbetta et al. (2008) that there should be a strong functional relation between the ventral frontoparietal network, involved in...
Commentary/Mather et al.: Norepinephrine ignites local hotspots of neuronal excitation

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Abstract: Emotional events can either impair or enhance memory for immediately preceding items. The GANE model explains this bidirectional effect as a glutamate “priority” signal that modulates noradrenaline release depending on arousal state. We argue for an alternative explanation: that priority itself evokes phasic noradrenaline release. Thus, contrasting E-1 memory effects are explained by a mechanism based on the Bienenstock–Cooper–Munro theory.

An emotional stimulus is typically well remembered but also influences memory for temporally adjacent events. In humans, we reported an emotion-induced retrograde impairment of memory in the context of shallow encoding of word lists containing an occasional emotional (E) noun (Strange et al. 2003). This retrograde disruption for “E-1” nouns appears to be mediated by the amygdala via a noradrenergic (NE) mechanism, as it is blocked by the β-adrenergic antagonist propranolol (Strange et al. 2003). However, subsequent studies have indicated that, if the encoding task requires that attentional weight be given to each E-1 stimulus, these stimuli show memory enhancement (Anderson et al. 2006; Knight & Mather 2009). Mather et al. propose that, for tasks involving attention to E-1 items, this “priority” signal is mediated by glutamate. According to their model, in a state of arousal, this elevated glutamate level associated with highly active neural representations stimulates greater NE release, leading to enhanced encoding of E-1 stimuli.

We propose that the opposing retrograde effects of emotion on memory can be explained by an alternative, simpler model. We propose that “priority” itself is coded by phasic NE release in the brain. Attending to task-relevant cues has been found to increase activity in the locus coeruleus (LC) in non-human primates (Aston-Jones et al. 1994). Thus, high “priority” E-1 encoding is likely to be associated with moderate levels of LC activity (Figure 1A, bottom). Given that enhanced memory for emotional items is blocked by propranolol, we assume that these emotional items provoke LC activity (Figure 1A, bottom). Because of the aversive nature of the E stimuli, this LC activity is likely to be greater than that evoked by task-relevant E-1 items. By contrast, in the case of low “priority” E-1 encoding, E-1 items trigger minimal LC activity (Figure 1A, top).

The bidirectional effects of emotion on memory for E-1 items can then be explained by a non-linear relationship between LC activity to E-1 items and memory encoding. According to the Bienenstock–Cooper–Munro model (Bienenstock et al. 1982),...
when the postsynaptic cell is weakly depolarized by other inputs, active synapses undergo long-term depression (LTD) as opposed to long-term potentiation (LTP) (Dudek & Bear 1992; Abraham & Tate 1997). The modification threshold, \( \theta_m \), is the measure of postsynaptic activity that determines the direction of synaptic efficacy change. In this scheme, if postsynaptic activity is below \( \theta_m \), but above baseline, synaptic efficacies are weakened. Conversely, if postsynaptic activity exceeds \( \theta_m \), synapses are strengthened. In Figure 1B, we apply this model to E-1 memory encoding (red curve). For low-priority E-1 items, postsynaptic activity is below \( \theta_m \) at the time of LC responses to the E noun, leading to a weakening of the efficacy of synapses that were engaged during the immediately preceding E-1 encoding (Diamond et al. 2004). For high-priority E-1 items, postsynaptic activity is already relatively high (above \( \theta_m \)) when the E stimulus is presented, yielding memory enhancement (red curve in Figure 1B). Note that the bidirectionality of this proposed effect is dependent on the presentation of E items. The black curve in Figure 1B represents memory for a stimulus that precedes a neutral (N) item (i.e., an N-1 stimulus) plotted as a function of the LC activity to this stimulus. Obviously, if, for any reason, this “N-1” stimulus evokes LC activity, its memory will be enhanced, but not to the level of enhanced E-1 memory. Importantly, N-1 memory will not be impaired even if it is low priority.

Thus, applying a model of the bidirectional nature of synaptic plasticity (Bienenstock et al. 1982) that has been validated in the context of NE stimulation (Hu et al. 2007; Kemp & Manahan-Vaughan 2008) can fully explain retrograde memory effects of emotion in a parsimonious way. The change in synaptic efficacy most likely occurs within a limited brain circuit involving amygdala and hippocampus (Strange & Dolan 2004), with NE input from the LC. It will be interesting to test whether contexts proposed to modulate \( \theta_m \), such as stress (Kim & Yoon 1998), will alter the direction of memory modulation for E-1 items for a given encoding task. Interestingly, blocking β-adrenergic receptors with propranolol does not abolish the emotion-induced retrograde amnesia for low-priority E-1 stimuli, but actually enhances memory for these E-1 items (Strange et al. 2003). It is tempting to speculate that propranolol decreases \( \theta_m \) (i.e., shifts the red curve in Figure 1b to the left), such that low levels of LC activity to low-priority E-1 nouns become associated with better memory.

Emotionally arousing context modulates the ERP correlates of neutral picture processing: An ERP test of the GANE model

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Abstract: The time scale of the effects of emotional arousal on neutral information processing is crucial for the predictions of the glutamate amplifies noradrenergic effects (GANE) model. GANE suggests that when emotional and neutral stimuli are presented in a sequence, neutral information processing will change. We review the literature on event-related potentials, including our own data set, to test this prediction.

The time scale of the facilitating versus impairing effects of emotional arousal on the processing of neutral information is an open question for the glutamate amplifies noradrenergic effects (GANE) model (sect. 2.1). The authors assert that “an emotionally salient word that impairs perception of a subsequent target word flashed in the same location 50 or 100 ms later can instead enhance perception of a target word flashed 1,000 ms later” (sect. 2.1). The specific time scales are likely to vary across experimental setups: for example, depending on the complexity of stimuli and the intensity of the arousal. The problem is that if the impact of arousal is not temporally bound, priority can be used to explain experimental effects in either direction, namely, both the impairing and the facilitating effects of arousal. Here we discuss how EEG data can provide crucial temporal dynamic information that can distinguish GANE’s predictions—evidence that Mather et al. did not consider.

According to GANE, arousing stimuli capture resources during their processing. Once their own processing is completed, the arousal they induce also facilitates the processing of subsequent stimuli. To test GANE we need to know in advance the duration of emotional stimulus processing. Previous work indicates that emotional pictures, a stimulus of choice in much of the human emotion–cognition literature, enhance a number of event-related potentials (ERPs). The most robust is the late positive potential (LPP). The LPP is thought to reflect attention allocation and maintenance of stimuli in working memory (Donchin & Coles 1988). The amplitude of the LPP 400–700 ms after stimulus presentation is higher when stimuli are emotional (Schupp et al. 2006), reflecting the additional resources allocated to such stimuli compared with GANE. Emotion also enhances other components, including the positive slow wave, where amplitudes are higher up to 6 s post-stimulus. We can therefore conclude that the processing of neutral information presented within 6 s of emotional pictures may be attenuated.

Only with this temporal information can we put GANE to the test. We do so by comparing the ERPs associated with processing neutral stimuli presented on their own (blocked neutral condition) with those presented alongside emotional stimuli (mixed condition). Not only is the context more arousing in the mixed condition (Long et al. 2015), but also emotional stimuli increase arousal locally. When the interstimulus interval (ISI) is long, emotional and neutral stimuli are unlikely to compete for processing resources, and GANE predicts that the higher global arousal in the mixed condition should enhance neutral information processing. In contrast, Pastor et al. (2008) used an ISI of 12 s and observed reduced LPPs for neutral stimuli compared with the blocked condition. It is, however, possible that those emotional pictures were still being processed when the subsequent picture was displayed after 12 s. When the ISI is short, competition should be pronounced, so GANE predicts that the processing of neutral information should be impaired. In contrast, Schupp et al. (2012) used an interval of 3 s and observed a null effect of context (blocked/mixed). It is, however, possible that the effect of emotion on resource allocation in that study was short-lived, for example, because of the orienting task. If emotional stimuli no longer attract attention when subsequent neutral stimuli are presented, the null effect is incompatible with GANE’s predictions.

In our experiment (Barnacle et al. 2015), 22 healthy adults viewed 16 lists of 14 pictures: 4 neutral, 4 emotional, and 8 mixed lists (50% emotional pictures). All pictures depicted people, emotional and neutral pictures. All pictures were equally semantically related, but the emotional pictures were more negative and arousing. Each picture was presented for 2 s with a jittered ISI of 4 ± 0.5 s. Participants were asked to encode these pictures for a free-recall memory test, which followed each study list after a 60-second distractor task. EEG was recorded during encoding with a BioSemi Active Two (BioSemi, Amsterdam) using 64 electrodes conforming to the 10–20 system and preprocessed with SPM (www.fil.ion.ucl.ac.uk). Data were filtered between 0.1 and 25 Hz, downsampled to 125 Hz, and epoched between ±200 and 4,000 ms. Eye-blink artifact was removed using an algorithm implemented in SPM. A threshold of 120 µV was used for trial rejection followed by robust averaging.

Following Schupp et al. (2012), we extracted LPP and slow-wave component amplitude data, averaging across centro-parietal electrodes (Cz, CPz, Pz, C1, C2, P1, P2, CP1, CP2) in three time windows: 400–700 ms, 1,000–2,000 ms, and 2,000–3,000.
ms poststimulus. We compared emotional and neutral picture processing in the mixed condition at each window to ascertain the duration of the effects of arousal, using three one-tailed paired-sample t-tests (p<0.017 controlled for multiple comparisons). Emotional modulated ERPs in the 400–700 ms and 1,000–2,000 ms windows, but not later (Figure 1). We then compared neutral picture processing in the mixed and blocked conditions at both these windows with two two-tailed t-tests (p<0.025). The LPP for neutral pictures was slightly, but not significantly attenuated in the mixed, compared with the blocked condition.

Our data indicate that the duration of the effect of arousal is key for testing GANE in novel experimental settings. EEG data allowed us to determine how long emotional pictures attracted extra processing resources. Here the modulation lasted up to 2 s from stimulus onset, evident in the modulation of the early portion of the LPP and positive slow wave, but not later. This pattern suggests that arousing stimuli are no longer in competition for resources when neutral pictures are presented in the same sequence. 3.5–4.5 s after the onset of the arousing pictures, and their prioritized processing should not detract from the resources allocated to neutral stimuli. On the contrary, because of the increased global arousal in the mixed condition, GANE predicts that neutral picture processing should be enhanced. In fact, our data provided evidence that the processing of neutral information is attenuated in that situation.

The three ERP data sets we reviewed appear to contradict GANE’s predictions. The mature electrophysiology literature on the effect of emotion on perception, attention, and memory (e.g. Hajcak et al. 2010; Schupp et al. 2006) can provide crucial evidence that the processing of neutral information is attenuated in that situation. However, a key focus of the BANE model was individual differences in prioritization of affectively salient stimuli (Markovic et al. 2014; Todd et al. 2012). This emphasis on individual differences has been based in part on observations of human carriers of a common deletion variant in the ADRA2b gene, which codes for noradrenergic α2B-autoreceptors (Small et al. 2001). The GANE model makes a valuable contribution in extending beyond the BANE model to incorporate the role of glutamatergic activity in enhancing effects of arousal on processing stimuli that are already high priority. However, the authors are somewhat dismissive of studies examining polymorphisms in genes coding for noradrenergic receptors and, specifically, of the notion that finding concerning the role of ADRA2b can be discussed in relation to the GANE model’s hotspot mechanisms. They do so based partly on evidence that α2B-receptors are unlikely to play an important role in GANE hotspots because the inhibitory role of α2B-receptors is not as well established as for α2A-receptors and because α2B-receptors are poorly expressed in key regions mediating affective salience. We argue that the study of genetic influences on affective prioritization of salient stimuli can provide data relevant to some of the GANE model’s claims, and that evidence against an inhibitory role of α2B-receptors in key brain regions is not entirely straightforward.

First, we argue that genotyping studies have value, in general, for understanding mechanisms of stimulus prioritization because, along with pharmacologic manipulations (e.g., De Martino et al. 2008; Strange et al. 2003), they are among the few vehicles for examining effects of inhibitory versus excitatory noradrenergic processes in humans. Because the specificity of ligands for receptor subtypes is limited (Jasper et al. 1998), genotyping studies can help specify the role of each subtype in patterns of brain activation and behavior. Of course, we acknowledge that it is important to use other methods, such as positron emission tomography and examination of mRNA activity, to help confirm the role of specific ADRA2a and ADRA2b polymorphisms in α2 activity.

Second, it is important to consider potential individual differences in the activity of specific receptor subtypes in proposed hotspot
processes and what the behavioral consequences might be. Genetic differences influencing such receptor function are one of such differences, and can provide a valuable window into how GANE mechanisms can vary normally and go awry. For example, common variants in genes coding for both ADRA2B and ADRA2A receptors have been associated with neural and behavioral indices of enhanced attention and memory for affectively salient stimuli that characterize affective disorders as well as cognitive biases associated with addictive behaviors (de Quervain et al. 2007; Havranek et al. 2015; Todd et al. 2013). Using genotyping to infer the role of each receptor subtype on such endophenotypes can help elucidate how patterns of inhibitory/excitatory activity proposed by GANE may contribute to variation in healthy populations and in psychopathology.

Studies of the ADRA2b deletion variant can serve precisely that function. Convergent evidence is highly consistent with the view that ADRA2b deletion carriers have reduced inhibitory autoreceptor function. In vivo, consequences of carrying the ADRA2b deletion variant (found in ≈50% of the populations we have studied) are similar to those of the α2-antagonist yohimbine (de Quervain et al. 2007). This claim is supported by the reliability and robustness of effects of enhanced emotional biases in attention and memory, increased amygdala and ventromedial prefrontal activation for arousing stimuli, and differences in amygdala gray matter volume associated with carrying the deletion variant (de Quervain et al. 2007; Ehlers et al. 2015; Rasch et al. 2009; Todd et al. 2013; 2014; 2015). According to the GANE model, affectively salient stimuli are one category of prioritized stimulus whose encoding is enhanced by arousal. Here, the enhanced affective prioritization we have observed in deletion carriers could lead to intensified positive feedback loops at hotspots, although possibly only when stimuli are prioritized because of their pre-existing associations with arousal. Further, because outside of the lab there are likely to be a range of motivationally relevant goals, behavior of deletion carriers may be driven by affective or visual salience over more “top-down” goals relative to non-carriers. Finally, with regard to the authors’ claims that it is α2B-autoreceptors that carry the full burden of inhibitory function in the brain, we suggest that the picture is somewhat more complicated. There is evidence that, in addition to its pre-synaptic inhibitory function, α2A is the most commonly observed postsynaptic receptor in the prefrontal cortex (Arnst et al. 1996; U’Prichard et al. 1979). Indeed, some evidence suggests that increased post-synaptic α2A activity in the PFC may be associated with enhanced rather than reduced noradrenergic transmission (Ramos et al. 2006). Moreover, brain regions mediating heightened emotional sensitivity in deletion carriers show relatively high levels of ADRA2b expression (Havrylyez et al. 2012; Allen Human Brain Atlas: http://human.brain-map.org). Animal research points further toward the importance of α2A-receptors in emotional processing (Moricau & Sullivan 2004). This challenges the notion of a straightforward role for α2A-receptors as the only mediators of inhibitory activity suggested by the GANE model.

In summary, although we acknowledge that effects of the deletion variant may be mediated by factors other than proposed GANE hotspot mechanisms, the growing body of research on polymorphisms influencing both a α2A- and α2B-receptors poses both questions and challenges for the GANE model.

Cognitive control, dynamic salience, and the imperative toward computational accounts of neuromodulatory function

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Author’s Response

GANEing traction: The broad applicability of NE hotspots to diverse cognitive and arousal phenomena

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Response/Mather et al.: Norepinephrine ignites local hotspots of neuronal excitation

the task-irrelevant information dominates processing later in the trial when cognitive control has prioritized task-relevant information, or both. Similar model simulations incorporating the timing of NE-mediated processing enhancements are also necessary to confirm whether GANE can account for the differential pattern of arousal effects on memory for stimuli occurring before and after arousing events (Sakaki et al. 2014a).

In principle, GANE may be implemented in the form of a detailed biophysical model (e.g., Eckhoff et al. 2009; Wang 2002) that simulates the cascade of neurochemical events at the “NE hotspots” described by Mather and colleagues. This component of the model would need to interact with other biophysically realized components that sustain associated cognitive functions (decision-making, cognitive control, memory) and generate task behavior, and the model predictions will depend on the interactions between these component processes and their relative timing. However, the fidelity of biophysical detail in such a model will likely trade off with its ability to provide a unified explanation of the vast array of arousal-related behavioral effects reviewed in Mather et al.

An alternative, perhaps more feasible approach would be for Mather and colleagues to adopt a simplified computational model of NE function that captures the essential impact of NE–glutamate interactions on task performance, in a form that is computationally tractable and can therefore be leveraged to generate predictions based on GANE principles in a wide variety of behavioral contexts. Indeed, a class of connectionist models of NE function already exists that appears well suited to such a pursuit. In these models, NE modulation is implemented as a multiplicative change in the input-to-output function of a task processing unit – otherwise known as a change in “gain” – and produces the critical winner-take-more/loser-take-less effects that GANE attempts to account for (e.g., Eldar et al. 2013; Servan-Schreiber et al. 1990). These models have been successfully adapted to explain neuromodulatory effects on perception and memory in a wide variety of task contexts, including those that require the online recruitment of cognitive control. Moreover, the model components governing NE modulation can be implemented at multiple levels of abstraction, from single model parameters that are global and time invariant (Eldar et al. 2013; Servan-Schreiber et al. 1990), to fine-grained subnetworks that operate on biophysically realistic principles and afford precise control over timing (Gilzenrat et al. 2002; Nienwenhuis et al. 2005b; Usher et al. 1999). In our view, whether the research question of interest pertains to arousal/cognitive control interactions or otherwise, this type of broadly applicable, computationally tractable modeling framework will be necessary to generate and test precise predictions of the GANE model in the future.

R1. Arousal

A number of the commentaries raise questions regarding arousal.

R1.1. Nature of arousal

In our view, the LC–NE system is not the only brain system involved in a generalized arousal response (see Pfaff 2006...
### Table R1. General topics raised in commentaries.

<table>
<thead>
<tr>
<th>What elicits LC activity?</th>
<th>Higher levels of arousal associated with <strong>uncertainty</strong> may help new salient information gain priority via hotspot mechanisms, whereas lower levels of arousal may protect existing strong predictions from distracting information under conditions of high certainty (Nassar, Bruckner, &amp; Eppinger [Nassar et al.]). <strong>Prediction errors</strong> may trigger a phasic NE response that facilitates the selective updating of predictions in the prioritized manner outlined by GANE (Ferreira-Santos). <strong>Competition</strong> elicits arousal, which leads to an increase in theta and gamma oscillations that select and stabilize “winning” representations (Phaf).</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forms of priority</td>
<td>Fluently processed stimuli yield a stronger signal (or are more salient), and so GANE can explain how arousal amplifies responses to these stimuli (Carbon &amp; Allbrecht).</td>
</tr>
<tr>
<td>How does GANE operate in relation to specific aspects of brain function?</td>
<td>Commentators discussed dendritic integration (Larkum &amp; Phillips), relative timing of oscillatory patterns (Phaf), the role of the dentate gyrus in memory selection (Houghton), and genetic variations in the ADRA2B gene (Todd, Ehlers, &amp; Anderson [Todd et al.]).</td>
</tr>
<tr>
<td>Spatial extent of hotspots</td>
<td>Eldar, Cohen, &amp; Liv (Eldar et al.) recognize that in the GANE model, hotspots would be co-extant with distributed cortical representations, whereas Gaucher &amp; Edeline are expecting more spatially extensive loci. This difference in visualization highlights the need for tools to identify active hotspot elements. Immediate early genes may be useful in this regard.</td>
</tr>
<tr>
<td>What are the adaptive functions of the neural effects of NE?</td>
<td>GANE may be a general-purpose function that cuts across a variety of cognitive and behavioral states (Hull). Acetylcholine is likely to have hotspot properties different from those of NE, and so low NE and high acetylcholine during REM sleep may help explain lack of memory for dreams (Becchetti &amp; Amadeo).</td>
</tr>
<tr>
<td>Relevance of GANE in various domains</td>
<td><strong>Stress.</strong> Endocrine signals, in particular cortisol, work in tandem with NE to promote long-term adaptive changes and memories (Hurleman, Maier, &amp; Scheele [Hurleman et al.]). <strong>Sleep and memory.</strong> Acetylcholine is likely to have hotspot properties different from those of NE, and so low NE and high acetylcholine during REM sleep may help explain lack of memory for dreams (Becchetti &amp; Amadeo). <strong>Early development.</strong> The LC exhibits developmental changes during infancy and early development, and early life stress shapes glutamate and GABA responses in ways that should be considered in the GANE model (Geva). <strong>Responses to sexual stimuli.</strong> Contrary to expectations of posture showing approach/avoidance biases, people viewing either threatening or sexual stimuli exhibit a freezing-like reaction in which they are more immobile (Mouras). <strong>Emotion regulation.</strong> Arousal levels should influence the ability to alter behavioral responses (Hull). <strong>Appraisal theory.</strong> Stimuli that are relevant for individuals’ goals, needs, and values induce strong arousal and amygdala activity (Montagrin &amp; Sander).</td>
</tr>
<tr>
<td>Factors that should be addressed</td>
<td>Commentators pointed out that GANE requires further development to specify timing (Talmi &amp; Barnacle; Navarra &amp; Waterhouse: Warren, Murphy, &amp; Nieuwenhuis [Warren et al.]), address different effects in prefrontal cortex (Abdallah, Averill, Krystal, Southwick, &amp; Arnsten (Abdallah et al.)), examine context and individual differences in determining salience (Huntsinger &amp; Storbeck), address role of ( \alpha )-receptors (Navarra &amp; Waterhouse), and address how cardiac afferents influence how LC modulates cortical activity (Critchley &amp; Garfinkel).</td>
</tr>
<tr>
<td>Alternatives to GANE</td>
<td>Priority is coded by phasic NE release and so there is no need for glutamate to signal priority (Strange &amp; Galarza-Vallejo; see response in sect. R4.1). The amygdala is necessary for NE to enhance selective processing and memory consolidation of arousing stimuli (Roozendaal, Luyten, de Voogd, &amp; Hermans [Roozendaal et al.]; see response in sect. R4.2). The tag-and-capture model is better able than GANE hotspot mechanisms to explain the effects of arousal on memories for events that occurred minutes to hours before the arousing event (Ritchey, Murty, &amp; Dunsmoor [Ritchey et al.]; see response in sect. R5). Countering the target article’s argument that a “network reset” model could not account for enhanced memory for well-attended items seen before an arousing event, Bouret argued that such enhanced memories could be accounted for by network reset if the qualitative nature of the representation changed (see response in sect. R6).</td>
</tr>
</tbody>
</table>
for a review of arousal pathways in the brain), but its activation is a common theme that runs through all different modes of arousal. For instance, NE inputs to cells in the ventromedial hypothalamus are critical in initiating sexual arousal (Pfaff 2006; of relevance for Mouras’ commentary), whereas noradrenergic input to the amygdala is critical in enhancing memory for emotionally arousing stimuli (see Roosendaal et al.’s commentary and sect. R4.2 on the role of the amygdala).

What is arousal? At the most basic level, we have the contrast between sleep and wakefulness. NE is low during most sleep states (see Becchetti & Amadeo). Then, during wakefulness, physical activity increases NE (Carter et al. 2010). But in addition to these broad-scale changes, the arousal system is also exquisitely sensitive and can adapt rapidly to small changes in the environment or internal goals.

These arousal responses can be detected by measuring pupil dilation. NE system activity increases pupil dilation, as NE released by the LC inhibits pupil constriction (Koss et al. 1984; Wilhelm 2008). Pupils are constricted during sleep, compared with wakefulness (Yoss et al. 1970). During wakefulness, aerobic exercise (Ishigaki et al. 1991) or muscular exertion (Nielsen & Mather 2013; Nielsen et al. 2015) increases pupil dilation. Arousal induced by stimuli or tasks also increases pupil dilation. For example, emotionally arousing scenes (Bradley et al. 2015), sexually arousing stimuli (Bradley et al. 2015), surprise, uncertainty, loud noises, and cognitive effort all increase pupil dilation. Subjective arousal ratings given for emotional images correlate with pupil diameter during viewing (Bradley et al. 2008). These consistencies across different elicitors of arousal suggest they share some underlying mechanisms to modulate cognitive and brain processing. Eldar et al. review a recent line of work in which they used pupil dilation as a marker of NE activity and found that indices of high NE function are associated with increased selectivity in learning, perception, and memory, consistent with their neural network models in which NE was modeled as global increase in gain. GANE complements and extends this approach by providing hypotheses about how NE implements neural gain.

We agree with Mouras and Kasper regarding the relevance of sexual arousal and internal sources of arousal (such as from one’s thoughts). Our point of view is that these different types and sources of arousal can be accommodated by the GANE model, as evidence suggests that LC activation is a common theme for all of them.

R1.3. How arousal may amplify the salience of negative stimuli

Kasper makes the case that negative stimuli may be more likely than positive stimuli to ignite neuronal hotspots because of the evolutionary pressure not to miss potential threats. One challenge is how to test this hypothesis, as negative stimuli, on average, induce more arousal than positive stimuli (Grühn & Scheibe 2008), and so any differences in processing or memory between negative and positive stimuli could be due to different levels of arousal when processing them, rather than to different levels of priority. To try to address this question, we recently ran a study in which we induced arousal independently by having participants squeeze a ball in their hand as hard as they could before they viewed emotional pictures and examined how the resulting increases in arousal influenced memory for the pictures (Nielsen et al. 2015). We were interested in hormonal effects, and all participants were younger female women. Consistent with Kasper’s predictions, we found that handgrip-induced arousal enhanced memory for the negative, but not the positive pictures. This effect was most pronounced for women with low estrogen and progesterone levels at the time of testing.

Kasper also suggested that because of declines in the LC–NE system, negative stimuli lose their arousing potential as people age. However, the evidence suggests that the older adults’ positivity effect is not due to a lack of bottom-
up salience for negative stimuli. Like younger adults, older adults look first at arousing stimuli regardless of their valence (Knight et al. 2007) and notice arousing or threatening stimuli more quickly than other types of stimuli (Leclerc & Kensinger 2008; Mather & Knight 2006). Bottom-up affective salience should play less of a role in influencing processing for low-arousal pictures, and indeed, the positivity effect appears to be stronger among valenced stimuli low rather than high in arousal (Kensinger 2008). In addition, we found that arousal induced by handgrip selectively benefited memory encoding of negative pictures (compared with positive or neutral pictures) in older women not taking hormone supplements, as well as in younger women with low estrogen and progesterone levels (Nielsen et al., in preparation). The evidence thus suggests that arousing negative pictures have similar bottom-up salience for older and younger adults.

R1.4. Relation between arousal and appraisal theory

On the basis of appraisal theory, Montagrin & Sander raise a question about how arousal and priority interact. They argue that arousal and goal relevance are not independent and stimuli that are relevant for individuals’ goals, needs, and values induce strong arousal and amygdala activity. We agree with them: Given that the LC exhibits phasic activity in response to goal-relevant stimuli (Aston-Jones & Cohen 2005; Aston-Jones et al. 1999), it seems possible that goal-relevant stimuli become arousing. However, the appraisal theory approach they discuss does not detail the neural mechanisms by which arousal induced by goal-relevant stimuli helps people memorize (Montagrin et al. 2013) and prioritize attention to those stimuli (Pool et al. 2015). In contrast, our GANE model can explain their findings of enhanced processing of goal-relevant stimuli: once the amygdala and/or prefrontal regions detect goal-relevant stimuli and recruit the LC (see Sara & Bouret 2012 for discussion of amygdala and prefrontal inputs to LC), NE hotspots will be generated in circuits transmitting goal-relevant information and, in turn, hotspots will enhance memory and perception for those stimuli. Therefore, GANE does not contradict the appraisal model, but instead extends it.

R1.5. Arousal and emotion regulation

Hull argues that the role of arousal in GANE is relevant for understanding impairments in emotion regulation. In particular, when stuck on a particular representation associated with negative emotions, decreases in arousal may be necessary to allow for less emotionally disturbing representations to be prioritized. Although not addressed in Hull’s commentary, a related point is the relevance of GANE for disorders such as post-traumatic stress disorder (PTSD), in which intrusive thoughts are a problem. A particular disturbing thought or memory may induce arousal, which, in turn, enhances attention to and memory reconsolidation of that particular representation. On the basis of GANE, beta blockers during initial encoding or retrieval of the memory should attenuate the immediate strength of its activation and its long-term synaptic strength. Consistent with this are some observational findings suggesting that beta blockers may help prevent intrusive thoughts or PTSD (Krauseneck et al. 2010; Lindgren et al. 2013), although random assignment has yielded some null effects (Stein et al. 2007).

R2. Priority

Other commentaries focused on physiological and psychological aspects of priority, a key factor in GANE.

R2.1. Perspectives on physiological mechanisms of priority

Larkum & Phillips describe a novel physiological mechanism by which contextual information modulates pyramidal cell activity. Neocortical pyramidal cell bodies have an apical trunk that ascends to a dendritic branching pattern called an apical tuft, which resides in a different cortical layer than the cell body and the basal dendrites around it. The long distance of the apical tuft from the cell body sets it up to serve a modulatory role in driving cell activity (Phillips 2015). Apical amplification could, for example, provide top-down priority selection of a quiet bottom-up auditory input to cortical output circuits. In their figure, Larkum & Phillips illustrate the interaction between GANE and apical amplification priority, providing an experimentally testable physiological model. Houghton argues that, computationally, the mossy cell hilar circuit in hippocampus would set priority for hippocampal processing and suggests heavy hilar NE innervation is consistent with GANE amplification of that mechanism. Becchetti & Amadeo make the interesting point that conscious (and, thus, prioritized) oneiric processing occurs during rapid eye movement (REM) sleep, likely supported by high acetylcholine modulation. But with active suppression of LC–NE during REM, there is little or no memory of those priority events, also consistent with GANE.

R2.2. Possible relation between fluency and priority

Carbon & Albrecht point out that fluency (i.e., processing information more easily) is an important factor determining stimulus priority. Greater fluency can arise because of perceptual salience (e.g., reading a word printed in a clear, high-contrast font more quickly than a blurry word) or because of prior knowledge or experience (e.g., reading a familiar word more easily than an unfamiliar word). Previous findings had suggested that people feel more positively about stimuli that they process more fluently (e.g., Winkleman & Cacioppo 2001). In a recent study, Albrecht and Carbon (2014) presented affective pictures that were either preceded (507 ms earlier) by that same image or by a different image shown for only 7 ms and asked participants to rate the valence of the pictures. There was no main effect of valence, but, instead, an amplification effect, with highly positive pictures rated more positively when they had been primed and highly negative pictures rated more negatively when they had been primed. Insomuch as fluently processed stimuli yield higher glutamatergic activity than less fluently processed stimuli (something that seems plausible but remains to be tested) and that the emotional stimuli elicited arousal, their findings that valence judgments of emotional stimuli are amplified by fluency fit with GANE.
R3. Predictive utility of GANE

Huntsinger & Storbeck and Talmi & Barnacle argued that GANE does not provide clear predictions concerning whether the presentation of emotionally arousing stimuli would enhance or impair cognitive processing of stimuli that appear nearby in time or space. Huntsinger & Storbeck state that GANE can provide post hoc explanations about the effects of emotional stimuli in a range of situations, but they question GANE’s predictive utility. Talmi & Barnacle also argue that because we don’t know exactly how long emotional stimuli dominate competition for representation, we can explain either the enhanced or impaired effects of emotional stimuli on nearby neutral stimuli by GANE.

We agree with them that it is hard to determine priority when comparing emotional with neutral stimuli. As discussed in our target article, emotional stimuli tend to have higher priority than neutral stimuli because of their goal relevance, bottom-up salience, and emotional salience. Thus, in the hypothetical experiment Huntsinger & Storbeck mention, where emotional stimuli are presented as distractors with task-relevant neutral stimuli, emotional distractors can have higher priority than neutral goal-relevant stimuli. This could especially be the case when the top-down control mechanisms are not strong enough to establish the goal relevance of neutral stimuli (see Warren et al.).

Talmi & Barnacle suggest that one can get around the issue of the different salience between emotional and neutral stimuli by having a long interval between emotional and subsequent neutral stimuli. But it is not clear that having a long interval would increase the priority of neutral stimuli as high as that of emotional stimuli. In addition, because high arousal can impair top-down prioritization (Arnsten 2011; Kulbhandan & Zehetleitner 2011), top-down control mechanisms might fail to increase the priority of neutral stimuli presented after emotional stimuli. These considerations suggest that in their EEG study (Barnacle et al., in preparation), neutral stimuli intermixed with emotional stimuli still had lower priority than neutral stimuli presented in a neutral list, which led to the impaired processing of neutral stimuli in the intermixed condition as predicted by GANE. Furthermore, having a long interval has the disadvantage that the effects of phasic arousal and NE release might not last long enough to yield modularly effects (see Section 9 in our target article).

In summary, it is difficult to test GANE in experimental settings where researchers simply include emotionally arousing stimuli and neutral stimuli without a clear manipulation of priority. In our view, to test GANE, it is important to manipulate the priority of neutral stimuli, independently from arousal (Lee et al. 2014b; Sakaki et al. 2014a; Sutherland & Mather 2012). One way to achieve this in the context of Barnacle et al. (in preparation) would be to have high-priority neutral images and low-priority neutral images in the mixed list condition. Similar changes can be made in the bridge study mentioned by Huntsinger & Storbeck (Dutton & Aron 1974); GANE predicts that arousal induced by the scary bridge will enhance memory for nearby high-priority stimuli (e.g., a woman seen on the bridge if the participant were asked to approach a woman and ask her something) while impairing memory for nearby low-priority stimuli (e.g., a man on the bridge who has no task relevance or particular interest). In summary, GANE can provide clear predictions as long as priority levels can be manipulated or assessed in the experiment.

R4. Alternatives to GANE proposed in commentaries

Several of the commentaries propose alternatives to GANE to explain the mechanisms by which arousing stimuli affect cognitive processing.

R4.1. NE-only model

Strange & Galarza-Vallejo propose that the glutamate aspect of the model is not necessary; they describe a simpler model in which priority is coded by phasic NE release in the brain. They work through an example from research on the emotional oddball–1 (E-1) effect, in which emotional oddballs (words or pictures) impair memory for the immediately preceding item on the list if that item was low priority for the participant, but enhance it if that item was high priority (e.g., Sakaki et al. 2014a). A problem with their NE-only model is that it is not clear how phasic NE release can selectively “tag” the E-1 item and not other items. Perhaps in the simple setup they describe, in which one word or object appears at a time in the list, phasic NE release could mark activated neural networks via a temporal tagging process. However, they do not consider findings that when multiple items are shown simultaneously, whether and how much memory for them is enhanced or impaired by a subsequent emotional item depends on their priority. For example, in an experiment in which a scene was shown either alone or with an object superimposed (Fig. R1A), if the image was followed by an emotional sound, there was impaired memory for the scene later, but only if it had been made lower priority by being in the background (Fig. R1B) (Ponzio & Mather 2014). Likewise, in another study in which participants saw four items at the same time that were then followed by a tone that was conditioned to predict either a shock (CS+) or no shock (CS–), having a subsequent arousing tone affected later memory for the simultaneously shown items differently depending on the relative priority of the items (Lee et al. 2015). The model Strange and Galarza-Vallejo propose does not explain how phasic LC activation could have different effects on items shown at the same time. In our view, this is the main contribution of GANE: by positing a mechanism for local cortical modulation of NE, it provides the only explanation to date of how arousal can have simultaneous differential effects on items based on their priority.

R4.2. Amygdala-based model

Roozendaal et al. argue that the amygdala is necessary for NE to enhance selective processing and memory consolidation of arousing stimuli. We agree that the amygdala plays a critical role, but argue below that its role in mediating the effects of NE is necessary only when the amygdala is the primary site of the neural representation in question.
Data from individuals with amygdala lesions help reveal which types of representations depend on the amygdala and which types can be supported by other brain regions. Compared with controls, unilateral amygdala patients exhibited as much enhanced visual cortex activity when viewing emotionally salient images (Edmiston et al. 2013), as much of an advantage for detecting emotional targets (Piech et al. 2010), and as much emotional capture by emotional stimuli during an attentional blink task (Piech et al. 2011). Two individuals with selective bilateral amygdala lesions exhibited a significant advantage in recalling aversive (compared with neutral) words during an attentional blink task, and this advantage was as large as that seen for matched control participants (Bach et al. 2011). Someone with complete bilateral amygdala lesions who could not recognize fear from faces still showed normal rapid detection of those faces (Tsuchiya et al. 2009). Thus, the amygdala is not necessary for the initial selective attention and encoding advantages seen for emotionally arousing stimuli, suggesting that NE–glutamate hotspots in sensory brain regions can occur even in the absence of the amygdala.

In addition, highly salient sensory stimuli yield normal physiological responses in people missing amygdalae (e.g., Tranel & Damasio 1989). For example, in studies of fear conditioning, individuals with amygdala lesions have normal skin conductance responses to aversive stimuli such as loud noises (Bechara et al. 1995; Klumpers et al. 2014). Likewise, three patients with bilateral amygdala lesions each had a panic attack when inhaling 35% CO₂ (Feinstein et al. 2013), indicating that amygdala lesion patients still experience fear in response to interoceptive alarming cues. These intact responses to interoceptive or external sensory stimuli contrast with the lack of fear shown by amygdala patients in response to experiences or visual stimuli (e.g., a haunted house or a live snake) that typically elicit fear because of their association with danger (Feinstein et al. 2011).

This pattern of findings suggests that the amygdala is essential for anticipatory physiological responses to stimuli that predict something aversive. This possibility is supported by fear conditioning studies with individuals with amygdala lesions (Bechara et al. 1995; Klumpers et al. 2014). These individuals lacked skin conductance responses to CS+ cues that predicted loud noises, even though they acquired explicit knowledge about the CS+ contingency. In contrast, an individual with bilateral hippocampal lesions failed to acquire explicit knowledge about the contingency, but had skin conductance responses to the CS+ (Bechara et al. 1995). Therefore, amygdala lesions impair physiological responding to cues that predict threat, but do not impair explicit learning about these cues. Amygdala lesions also impair physiological responding to cues that predict something positive or negative to yield a physiological affective response.

The findings that patients with amygdala lesions no longer have physiological responses to predictive cues...
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despite having as much explicit knowledge of the contingencies as normal controls suggests that: (1) there are amygdala-based neural representations of associations between neutral cues and potential affectively relevant outcomes; and (2) these amygdala-based representations are necessary to trigger signals to sympathetic pathways to mount a physiological response, possibly in part via amygdala projections to the LC (Cedarbaum & Aghajanian 1978).

Likewise, the finding that an individual with a hippocampal lesion lacked explicit knowledge of fear conditioning contingencies despite exhibiting a skin conductance response to the CS+ suggests that there also are amygdala-independent, hippocampus-based neural representations of associations between CSs and USs. However, in people with intact amygdalae and hippocampi, these separate representations in the two regions are likely to have close interactions, in part supported by a direct glutamatergic pathway from the basolateral amygdala to the CA1 region of the hippocampus (Rei et al. 2015).

Noradrenergic contributions to interactions between amygdala and hippocampus have been examined using one-trial learning to avoid a shock (McIntyre et al. 2005). In this paradigm, the β-adrenergic receptor agonist clenbuterol is infused into the basolateral complex of the amygdala shortly after a rat learns that moving from a brightly lit compartment of an alley through a door to a dark compartment is associated with a shock. The β-adrenergic stimulation of the amygdala increases Arc expression (indicating more synaptic plasticity) in the hippocampus in the 45 min after the shock. Of particular relevance in this context, however, are findings that the increased Arc expression depends not only on greater NE activity in the amygdala itself, but also on arousal levels more generally (McReynolds et al. 2014).

Specifically, whereas basolateral amygdala infusions of a β-agonist increased Arc protein levels for the inhibitory avoidance shock task, as seen in previous studies and also for a “high-arousal” version of an object recognition task, NE activity in the amygdala was not sufficient to increase Arc in the hippocampus when the object recognition task was not arousing. These findings suggest that glutamate–NE feedback loops in the amygdala can be intensified by within-amygdala local β-adrenergic activation (Fig. R2A). This hotspot activity increases glutamatergic signaling to the hippocampus (Fig. R2B) but does not directly increase NE levels in the hippocampus. However, the increased glutamatergic activity in the hippocampus can stimulate local release of NE via NMDA receptor activity at LC neuron varicosities if the LC is depolarized (Fig. R2C; see target article for more details on hotspot mechanisms). In summary, McReynolds and colleagues’ data suggest that NE can influence hippocampal activity either indirectly via glutamatergic pathways from the amygdala or directly via local release from LC varicosities. More generally, we posit that NE action within the amygdala has important glutamatergic modulatory effects elsewhere in the brain (in particular in the hippocampus), but that the LC also modulates excitation and inhibition directly in these other brain regions via local release of NE. The critical experiments necessary to test this hypothesis have not been performed yet (see relevant proposed study in Table R2).

Roozendaal et al. also argue that “the impairing effects of amygdala–NE interactions on memory of non-salient/non-arousing information involve an active process that is dependent on the amygdala.” They make this case based on Lovitz and Thompson (2015), whom they interpret as showing that intra-basolateral amygdala infusion of a β-adrenergic agonist (clenbuterol) decreases hippocampal excitability in non-inhibitory avoidance-trained control animals. However, their interpretation appears to be incorrect, as in that study, there was no significant difference between vehicle and clenbuterol conditions in the untrained rats.

R5. Role of NE hotspots in long-term memory formation

Some commentators raise questions concerning the role of NE hotspots in memory. First, Hurlemann et al. point out the importance of cortisol, in addition to NE and glutamate, in explaining the effects of arousal on memory. Combining neuroimaging with a psychopharmacological approach, Hurlemann et al. demonstrated that NE and glucocorticoids interact during processing of emotional stimuli (Hurlemann 2008; Kukolja et al. 2008, 2011). In particular, their work suggests that NE interacts with cortisol to enhance learning of emotional information within the amygdala–hippocampus network.

Acute stress and administration of glucocorticoids lead to enhanced glutamate release both in the amygdala (Reznikov et al. 2007) and in the hippocampus (Moghaddam et al. 1994) via mechanisms mediated by glucocorticoid receptors (GRs) and mineralocorticoid receptors (MRs) (for reviews, see Popoli et al. 2012; Sandi 2011). In the amygdala and hippocampus, interactions between glucocorticoids and NE have been observed, as well (for reviews Joels et al. 2011; Krugers et al. 2012). These results suggest the interesting possibility that glucocorticoids help NE create hotspots in the amygdala–hippocampus circuit by enhancing glutamatergic activity. One question is whether the NE–cortisol interaction goes beyond the amygdala–hippocampus circuit. Although most previous research focuses on the effects of glucocorticoids either in the amygdala-hippocampus pathway or in the prefrontal cortex (PFC), glucocorticoids might also amplify NE hotspots in other cortical regions, given that GRs are widely expressed in brain (Morimoto et al. 1996). Furthermore, elevated cortisol and NE levels tend to impair goal-directed attentional processes in the PFC (Schwabe et al. 2012), which should enhance the impact of the bottom-up, salience-driven hotspots predominant in sensory brain regions.

Second, Ritchey et al. state that the tag-and-capture model is better able than GANE hotspot mechanisms to explain the effects of arousal on memories for events that occurred minutes to hours before the arousing event. For example, initially weak memories can be strengthened by a subsequent salient signal, such as a novelty or aversive event (Dunsmoor et al. 2015; Redondo & Morris 2011). The tag-and-capture model explains these results by asserting that memory traces are tagged during initial learning, which allows for subsequent plasticity-related protein-mediated mechanisms to capture those tagged traces to create long-term memories. Ritchey et al. also argue that the effects of arousal on protein synthesis processes are mediated by dopaminergic neuromodulation.
Although in our target article we focused mainly on the immediate effects of NE hotspots, we believe that evidence indicates a role for these hotspots in tag-and-capture scenarios. \(\beta\)-Adrenergic receptor activity stimulates protein synthesis and gene expression alterations associated with long-term potentiation maintenance (Maity et al. 2016; O’Dell et al. 2010). NE hotspots should play a role in tag-and-capture by elevating local NE levels to activate \(\beta\)-adrenergic receptors, as well as by increasing glutamatergic activation of NMDA receptors. Both \(\beta\)-adrenergic activity and NMDA activity (in addition to dopamine D1/D5 receptor activity) are essential to “set the learning tag” for an initial weak memory, and \(\beta\)-adrenergic receptor activation is required during exposure to the modulating novel event occurring an hour later (Moncada et al. 2011). A particularly intriguing finding is that the behavioral tagging phenomenon requires the initial weak event and the subsequent novel event to occur in the same sensory modality, thereby activating the same general population of neurons (Ballarini et al. 2009). Likewise, Dinnsmore et al. (2015) found that fear conditioning enhanced memory for previously learned images only when those images were semantically related to a fear-conditioned category; when images of animals were fear-conditioned, memories for previously learned animals were enhanced, whereas when images of tools were fear-conditioned, memories for previously learned tools were enhanced. This is consistent with the local nature of NE hotspots and raises the

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**Fig. R2.** (Mather et al.) Previous findings suggest that NE can influence hippocampal activity both directly from NE release from LC neurons and also via amygdala glutamatergic pathways. (A) LC neurons innervate both amygdala and hippocampus. (B) NE released during LC activation (or, in the case of McReynolds et al., 2014 discussed in the text, a \(\beta\)-agonist) interacts with activated local glutamatergic representations within the amygdala to create a hotspot of higher glutamatergic activity. (C) These glutamate-NE hotspots originating in the amygdala amplify hippocampal glutamatergic activity via glutamatergic pathways. Neurons originating in the amygdala connecting to the hippocampus do not release NE, and so amygdala activity will not directly affect NE levels in the hippocampus. (D) However, amygdala modulation of hippocampal glutamatergic activity can increase the probability of local hotspots developing in the hippocampus, because if the LC is depolarized, the amygdala-induced glutamatergic activation in the hippocampus stimulates local NE release and further amplifies glutamatergic activation via glutamate-NE hotspot mechanisms. (E) Even in the absence of amygdala modulatory activity, NE hotspots can develop in the hippocampus from excited glutamatergic activity interacting with NMDA receptors on local LC neuron varicosities to increase local release of NE, as demonstrated in hippocampal slice preparations (e.g., Lalies et al., 1988; Nelson et al., 1980; Pittaluga & Kaiteri, 1990; Vezzani et al., 1987; Wang et al., 1992). Note: Inset in (A) is reprinted with permission from Marien et al. (2004), p. 41, part of Figure 2.
Table R2. Data needed to test hypotheses and better understand arousal–priority or NE–glutamate interactions.

<table>
<thead>
<tr>
<th>Question</th>
<th>Proposed Experiment/Method</th>
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<tbody>
<tr>
<td>Can we measure GANE-proposed neurotransmitter mechanisms in laboratory animals?</td>
<td>Direct measurements of local glutamate levels and NE or β-adrenergic receptor activation levels in awake cortex with arousal/cue manipulations would make it possible to test our physiological GANE model. New techniques make it possible to track extra-synaptic glutamate activity (Okubo et al., 2010), and researchers are getting closer to being able to monitor levels of NE and G-couple protein receptor activation at spatial resolutions corresponding to a representational network (Muller et al., 2014).</td>
</tr>
<tr>
<td>Does NE interact with apical amplification priority signaling?</td>
<td>The Larkum &amp; Phillips hypothesis that NE modulates apical amplification in the output neurons of cortex as the mediator of top-down or cortico-cortical priority signals can be examined both in vitro and in vivo. Evidence for such gating would significantly expand the GANE model.</td>
</tr>
<tr>
<td>Is “network reset” a general motor–sensory or structure-specific effect?</td>
<td>Immediate early genes with the ability to reveal two brain activation sequences separated by a temporal interval could test the reset (reorganizing)-versus-amplification effects of phasic LC activation. We predict evoked sensory representations would be enhanced and stabilized by phasic glutamatergic activation of LC, whereas hippocampal and possible prefrontal representations would be reconfigured. Tonic effects of NE would not evoke reset.</td>
</tr>
<tr>
<td>How close in time does phasic arousal need to be to modulate the priority of another event?</td>
<td>Initial behavioral data suggest that arousal induced by one event can modulate processing of other events occurring within a few seconds (see target article for review). Previous work indicates that glutamate activation of NMDA receptors decays slowly and can last hundreds of milliseconds (Lester et al., 1990), but more work is needed to quantify the timing of glutamate and NE actions at hotspots (allowing for formal modeling, as highlighted by Warren et al., in their commentary).</td>
</tr>
<tr>
<td>Can we measure GANE-proposed neurotransmitter mechanisms in humans?</td>
<td>Advances in human magnetic resonance spectroscopy (MRS) enable the measurement of glutamate metabolites in vivo, but with poor spatial and temporal resolution. One straightforward test of GANE would be to examine whether an arousing stimulus can elicit a local, activity-dependent increase in glutamate levels for a prioritized stimulus.</td>
</tr>
<tr>
<td>Test of NE hotspots in humans</td>
<td>During task-related fMRI involving an arousal/priority manipulation, trial-by-trial estimates of pupil dilation to the arousing stimulus could be used to scale BOLD responses in cortical representational regions underlying the high-priority stimulus. This would provide an estimate of how LC responses selectively modulate local cortical activity.</td>
</tr>
<tr>
<td>Test Roozendaal et al. argument that NE effects on memory rely on the amygdala.</td>
<td>The fact that the hippocampus has many NE receptors suggests that NE can modulate memory consolidation in the hippocampus directly, without amygdala modulation (although NE release in the amygdala can lead to glutamatergic activation of the hippocampus, it does not directly increase NE in the hippocampus; see Fig. R2). A simple experiment would be to attempt to modulate consolidation of a hippocampally represented memory such as learning the context of a novel object by infusing NE into the hippocampus (as has been done with NE infused into the amygdala [Barsegyan et al., 2014]).</td>
</tr>
<tr>
<td>Inverted-U curve</td>
<td>A direct examination of inverted-U curve effects with NE would be of interest. It is not clear if the functional shift seen at high levels of arousal is uniquely, or even critically, due to high NE levels or is a multifactorial effect depending on co-activation of other systems.</td>
</tr>
</tbody>
</table>
interesting question of just how widely the plasticity-related proteins stimulated via β-adrenergic receptor activation at NE hotspots modulate interconnected memory circuitries. The behavioral findings (Ballarini et al. 2009; Dunsmoor et al. 2015) suggest that they do not have an influence much beyond a local region that represents the same category or sensory modality of item. Although much still needs to be worked out about the potentially complementary roles of dopamine and norepinephrine on tag-and-capture phenomena, we believe that thinking about the local nature of the β-adrenergic activity induced by arousing modulatory events will be fruitful.

R6. GANE amplification of prioritized representations during a “network reset”

According to a prominent theory, NE release orchestrates a “network reset” that reorients attention and, consequently, reorganizes underlying representational networks during a sudden and unexpected change in environmental impertives (Bouret & Sara 2005; Sara & Bouret 2012). We agree with Sara’s perspective that GANE is complementary to the “reset” hypothesis. From the perspective of GANE, whether this type of reorienting occurs will depend on whether there are currently representations with high glutamatergic activity or not. If there are no current strongly active representations, both GANE and the network reset theory predict that the predominant effect of an increase in LC activity would be to enhance reorienation to new salient stimuli. However, when there is already a highly active representation, GANE predicts that an increase in LC activity will further enhance processing of that representation (e.g., Anderson et al. 2006; Knight & Mather 2009; Sakaki et al. 2014a), rather than having a network reset effect. On the basis of these findings, in our target article we argued that the network reset perspective fails to account for the ability of arousal to enhance memory of preceding high-priority information. Bouret responded by suggesting that enhanced memory for a preceding event could be consistent with a network reset if, when an arousing event occurred, the preceding salient event was now represented in a qualitatively different way that was integrated with the arousing event.

Consistent with Bouret’s argument that arousal enhances memory for preceding information when the preceding information is integrated with the arousing events, in fear/evaluative conditioning paradigms events repeatedly followed by emotional outcomes acquire emotional properties (for a review, see Baeyens et al. 2005). Our previous research also demonstrated that when individuals are presented with neutral cues followed by emotional or neutral outcomes, emotional outcomes facilitate memory for neutral cues only when they are aware of the cue-outcome contingency (Mather & Knight 2008; Sakaki et al. 2014b).

To address the important question raised by Bouret about whether arousal changes the nature of representations, future research should probe the effects of arousal on the specificity of mental and neuronal representations. At least one recent study suggests active sensory representations are strengthened, rather than altered, by noradrenergic system activation (Shakhawat et al. 2015). In addition, our findings suggest that emotional arousal enhances the veracity of the original representation, or detail memory, rather than gist alone (Sakaki et al. 2014a).

R7. Alternative ways to trigger LC activity

Although most of the target article focused on how emotionally arousing stimuli shape cognitive processing, non-emotional stimuli can also activate the LC and thereby influence cognition. In this section, we discuss how prediction errors, uncertainty, and competition each influence LC activity.

R7.1. Prediction errors

Prediction is a central feature of efficient cognitive processing. As described by Ferreira-Santos, GANE fits well with “predictive coding” frameworks of cognition: Sudden mismatches between predicted and actual sensory and affective inputs represent an important form of conflict and competition that can elicit arousal and LC activity. Supporting this view, pupil dilation has been linked to the occurrence of prediction errors (Braem et al. 2015; Preuschoff et al. 2011). Furthermore, in monkeys, phasic LC activity ceases to signal the occurrence of reward once the reward follows a specific action predictably (Sara & Segal 1991). Other research also indicates that affect enhances prediction error responses (Vogel et al. 2015a) and that prediction errors are a fundamental component of generating interceptive feelings (Barrett & Simmons 2015).

R7.2. Uncertainty

As pointed out by Nassar et al., as well as by Bouret, it is important to consider the purpose of having one level of arousal modulate cognitive processing differently than another level. When is it useful for cognitive processing to remain focused on previously salient information? And when will it be advantageous to be open to new prioritized information? Nassar and colleagues argue that during times of uncertainty, it is especially important not simply to focus on current prioritized cues, but to amplify incoming prioritized sensory information (Yu & Dayan 2005). They review findings that pupil diameter is larger during periods of uncertainty than when expectations are reliable. Thus, tonically higher levels of NE should decrease the threshold for new salient stimuli to ignite hotspots. They suggest that older adults’ deficits in learning under conditions of uncertainty may be linked to age-related declines in LC function.

R7.3. Competition and conflict

As highlighted by Phaf, there is much evidence that competition and conflict between representations induce arousal. These stimuli/events are also likely to produce hotspots, based on evidence that conflict, along with novelty, target detection, uncertainty, and performance errors, elicit LC activity (for reviews, see Berridge & Waterhouse 2003; Nieuwenhuis et al. 2005a; Ullsperger et al. 2010; Yu & Dayan 2005). Fundamentally, GANE predicts that any stimulus that activates the LC–NE system will produce hotspots in an activity-dependent manner, regardless of
whether NE release is triggered by something emotional or not. If competition elicits arousal, it could very well be an effect driven by prediction errors (i.e., significant discrepancies between feedforward and feedback inputs; see sect. R7.1), initiating a network reset via the LC.

Phaf also discusses the distinct but complementary roles of theta and gamma oscillations in signaling and resolving stimulus conflict, respectively. According to Phaf, theta arises from conflict, is a substrate of arousal, and helps select dominant representations via intercortical communication. Subsequently, gamma oscillations facilitate a resetting and stabilization of “winning” representations. His description is consistent with Sara’s empirical data. In her commentary, Sara describes evidence that stimulating the LC briefly suppresses gamma oscillations for 200 ms, which is followed by a near doubling of the gamma power immediately afterward, as well as an increase in theta power (Sara 2015). Interestingly, in an early report of conflict activating LC, the absence of expected reward elicited a specific theta band increase (∼7.7 Hz) in hippocampus (Gray & Ball 1970). This effect was later demonstrated to require forebrain norepinephrine (Gray et al. 1975). It could be useful to reexamine this theta signature of LC activation (for more recent support, see Walling et al. 2011) and its role in synchronizing activity for prioritized representations. Another interesting question is whether (as suggested in the target article) NE hotspots enhance local gamma power via a β-adrenergic pathway, thereby increasing selective attention.

R8. Additional mechanistic considerations/complications for GANE

As noted by several commentators, GANE is necessarily a simplification of a complex reality. It does not, for example, incorporate the function of postsynaptic α2-receptors, the subthreshold input promoting role of α1-receptors, the synergistic role of α1- with β-adrenergic receptors or recently described astrocytic functions of α1-receptors. The co-release of peptides from LC varicosities is not considered; neither is the probable role of other neuromodulators known to be elevated in various forms of arousal discussed. This is a beginning that will, ideally, lead to a more veridical model of cortical self-regulation that addresses how neurotransmitters released during arousal interact with local cortical conditions to modulate activity in flexible yet highly targeted ways.

R8.1. Varied effects of adrenoreceptors

As highlighted in several commentaries, the GANE model does not incorporate all known adrenoreceptor functions. These omissions include the role of postsynaptic α2-receptors that play important roles in the PFC (see commentaries by Abdallah et al. and Todd et al. and that also occur in other areas of neocortex (Venkatesan et al. 1996). Navarra & Waterhouse and Gaucher & Edeline point out that α1-adrenoreceptors have more varied actions, including synergism with β-adrenoreceptor effects, potentiation of effects on their own, and astrocytic action. In particular, they highlight that the role of α2-adrenoreceptor in sensory cortex may be facilitatory. When activated, these receptors appear to potentiate postsynaptic excitatory responses and can boost subthreshold inputs (for a review, see Berridge & Waterhouse 2003). Furthermore, global astrocytic calcium waves are initiated via LC–NE activation of astrocytic α1-adrenoreceptors (Ding et al. 2013), consistent with a model in which LC–NE global effects recruit both α1- and α2-adrenoreceptors.

R8.2. Suppressive effects of NE in sensory regions

Gaucher & Edeline emphasize the suppressive actions of exogenous NE on processing in auditory cortex as being inconsistent with GANE. But their finding that a small population of auditory neurons encoding natural stimuli are enhanced by NE (Gaucher & Edeline 2015) and contribute to discrimination is similar to newer findings in olfactory cortex that LC–NE modulation is essential for difficult natural odor discrimination and increases the stability of small distributed odor representations (Shakhavat et al. 2015), as predicted by GANE.

R8.3. Differential effects of adrenergic receptors in prefrontal and posterior cortex

Abdallah et al. highlight the differences between the actions of NE on classic sensory synapses in subcortical and posterior sensory regions and newly evolved circuits in layer 3 of the dorsolateral PFC (DLPFC). On the basis of animal and human research, they suggest hotspot effects are most likely to occur in sensory and limbic (e.g., amygdala, hippocampus) synapses where β-adrenoreceptors promote glutamate responses and long-term potentiation. In the PFC, in contrast to “classic” sensory areas, β-adrenoreceptor activation has been found to impair rather than enhance postsynaptic function via increased cAMP signaling (Arnsten et al. 2015a; Ramos & Arnsten 2007). Like β-adrenoreceptors, α1- and α2-adrenoreceptors also appear to have contrasting influence on neuronal activity in the PFC versus sensory cortices. Although α1-receptors enhance sensory neuron firing, they tend to impair PFC function and working memory (Ramos & Arnsten 2007). On the other hand, whereas α2-receptors enhance inhibitory signals and suppress noisy activity in the posterior cortex, their activation strengthens dorsolateral PFC functional network connectivity and promotes working memory (Arnsten et al. 2012). These inverted rules of adrenoreceptor function in the PFC have important implications for how GANE influences cognitive processing during sudden arousal. Although an arousal-induced surge of NE may disrupt working memory representations in the DLPFC (e.g., current event models), it should also transiently enhance the throughput of strong glutamatergic signals in the hippocampus (Brown et al. 2005). Therefore, DLPFC impairments may facilitate reorientation during arousal to information that has bottom-up salience and is associated with hotspots of high activity in sensory regions but not in PFC.

R8.4. Relative timing of arousal and prioritization process

The key distinction outlined in the previous section between the effects of NE in sensory cortices and limbic regions versus the PFC agrees well with the timing hypotheses proposed by Warren et al. In their commentary,
Warren and colleagues present evidence that the relative strength of bottom-up and top-down (cognitive control) priority inputs changes rapidly within a single trial. Whereas bottom-up salience dominates the competition for mental resources early on, cognitive control processes take longer to develop and overcome the initial dominance of perceptual salience. Warren et al. suggest that this time-variant model of salience determines whether phasic arousal enhances or impairs task-relevant (but not perceptually salient) information.

Indeed, the GANE model predicts that arousal-induced NE release will bias competition in favor of whatever information has the highest priority at that moment. Experiencing arousal while a representation is highly active should strengthen that representation regardless of whether top-down goals or bottom-up salience prioritized the representation, because the representation was activated before moderate to high levels of NE could disrupt goal-directed processing in the PFC (Ramos & Arnsten 2007). In contrast, the source of priority may matter more when experiencing arousal before a stimulus is perceived. Although prestimulus arousal should amplify the effects of bottom-up salience, it may diminish the effects of top-down priority if, as outlined in the previous section, working memory processes that help maintain and implement processing goals are impaired by the arousal (Ramos & Arnsten 2007).

Data from our lab provide clear evidence that prestimulus arousal enhances the impact of bottom-up salience (Lee et al. 2014b; Sutherland & Mather 2012), whereas poststimulus arousal enhances the impact of top-down prioritization (Lee et al. 2015; Sakaki et al. 2014a). Whether arousal enhances priority for the other two combinations remains to be seen. We have not yet tested scenarios in which something perceptually salient is followed by something arousing, but GANE would predict that as long as the representation associated with that perceptually salient item were still strongly active when arousal increased, it might enhance rather than disrupt the representation associated with that perceptually salient thing arousing, but GANE would predict that as long as the representation was activated before moderate to high levels of NE could disrupt goal-directed processing in the PFC (Ramos & Arnsten 2007). In contrast, the source of priority may matter more when experiencing arousal before a stimulus is perceived. Although prestimulus arousal should amplify the effects of bottom-up salience, it may diminish the effects of top-down priority if, as outlined in the previous section, working memory processes that help maintain and implement processing goals are impaired by the arousal (Ramos & Arnsten 2007).

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8.6. Individual differences

Geva points out that tonic levels of arousal predict whether infants orient toward novel or familiar stimuli, and suggests that infancy is an interesting test case for GANE, as, unlike in later stages of development, infants lack an “established neural network set with implicit know-hows” that provide the glutamatergic priority signal necessary to ignite hotspots under arousal. Differences at the other end of life are also relevant, as Nassar et al. point out. Genetic variation in adrenergic receptors also may matter; Todd et al. make the case that ADRA2b deletion carriers have reduced inhibitory autoreceptor function.

9. Conclusions

As evinced by the diverse range of commentary, the NE hotspot mechanism goes beyond just the emotion–cognition literature to explain how arousal influences different forms of cognitive selectivity. One of GANE’s most vital contributions is that it showcases the ability of the cortex to regulate its own processing efficiency. Such local control of cognition represents a fundamental mechanism of adaptive brain function that has the potential to explain a variety of cognitive phenomena. As GANE exemplifies, synaptic activity is not just passively modified by neuromodulators. Instead, under situations of arousal that demand our attention, such as threat or excitement, salient brain signals recruit the ingredients necessary to form lasting memories.


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