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511-fear-stress

Rick Gilmore

2021-11-11 13:56:09

- Fun
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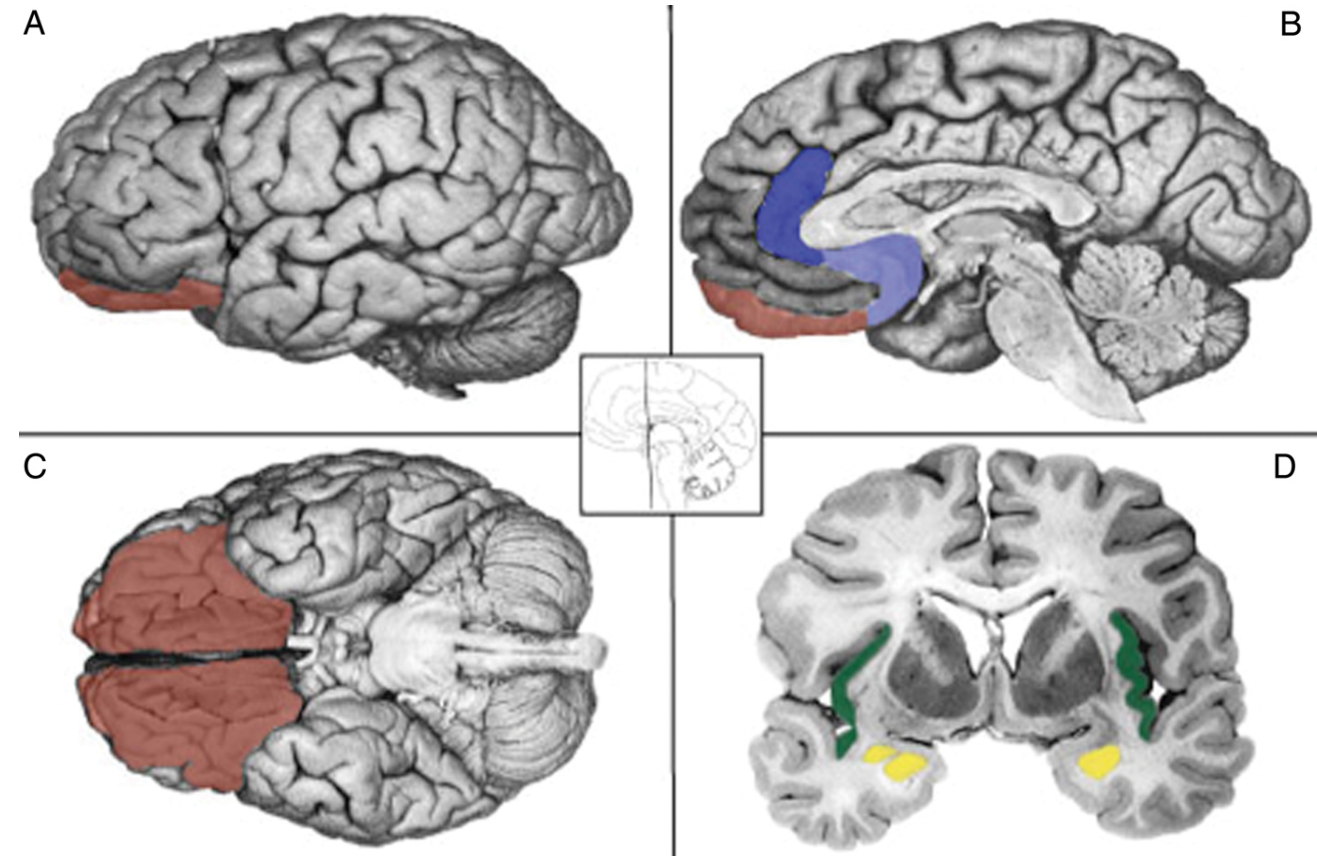
Fun

The brain bases of emotion

(Lindquist, Wager, Kober, Bliss-Moreau, & Barrett, 2012)
(<http://dx.doi.org/10.1017/S0140525X11000446>)

Locationist account

- *Where* in the brain is emotion processed?



(Lindquist, Wager, Kober, Bliss-Moreau, & Barrett, 2012)

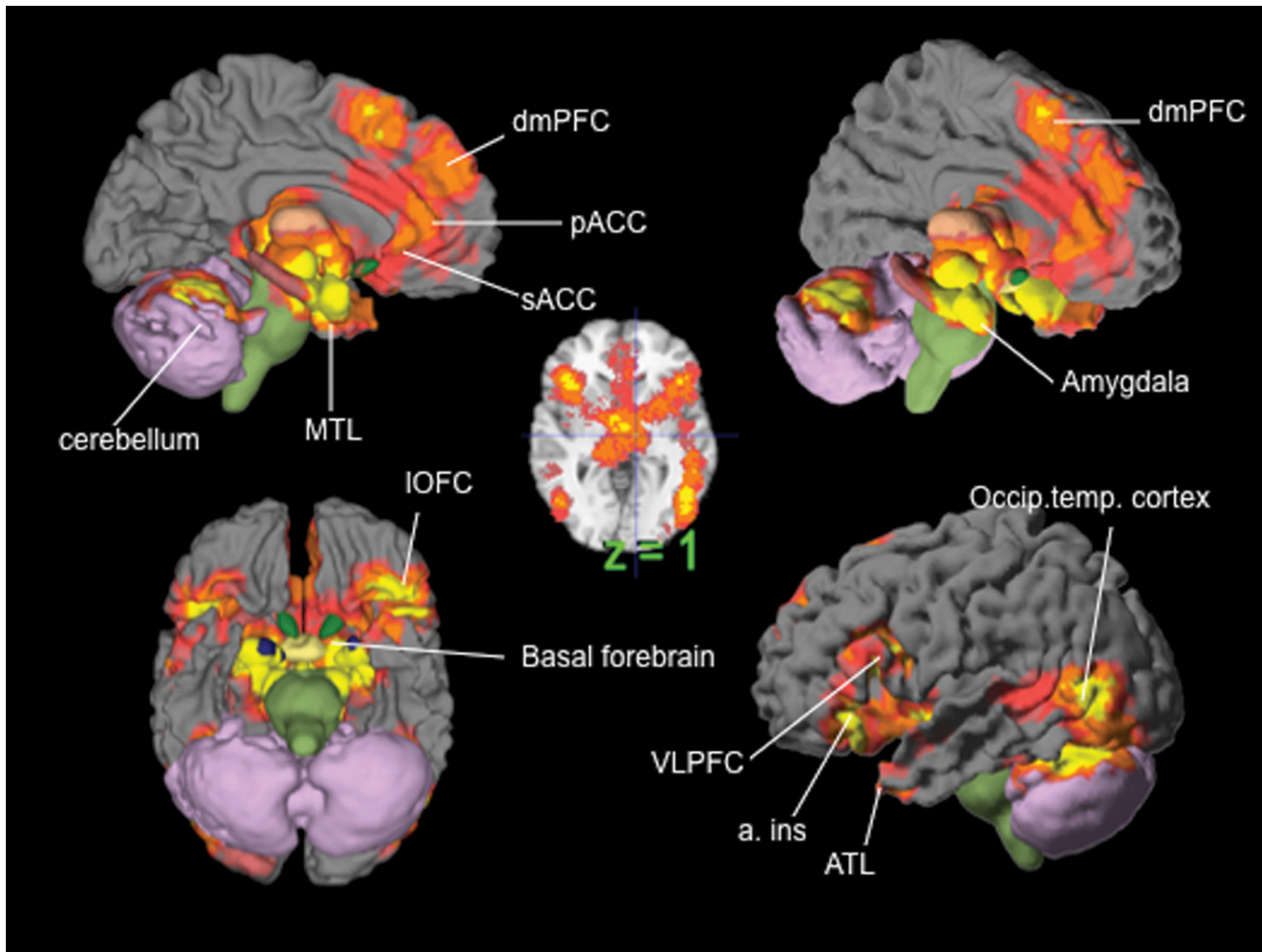
(<http://dx.doi.org/10.1017/S0140525X11000446>)

Figure 1. Locationist Hypotheses of Brain–Emotion Correspondence. A: Lateral view. B: Sagittal view at the midline. C: Ventral view. D: Coronal view. Brain regions hypothesized to be associated with emotion categories are depicted. Here we depict the most popular locationist hypotheses, although other locationist hypotheses of brain–emotion correspondence exist (e.g., Panksepp, Reference Panksepp1998). Fear: amygdala (yellow); Disgust: insula (green); Anger: OFC (rust); Sadness: ACC (blue). A color version of this image can be viewed in the online version of this target article at <http://www.journals.cambridge.org/bbs> (<http://www.journals.cambridge.org/bbs>). (Lindquist, Wager, Kober, Bliss-Moreau, & Barrett, 2012) (<http://dx.doi.org/10.1017/S0140525X11000446>)

Constructionist account

A psychological constructionist account of emotion assumes that emotions are psychological events that emerge out of more basic psychological operations that are not specific to emotion. In this view, mental categories such as anger, sadness, fear, et cetera, are not respected by the brain (nor are emotion, perception, or cognition, for that matter).

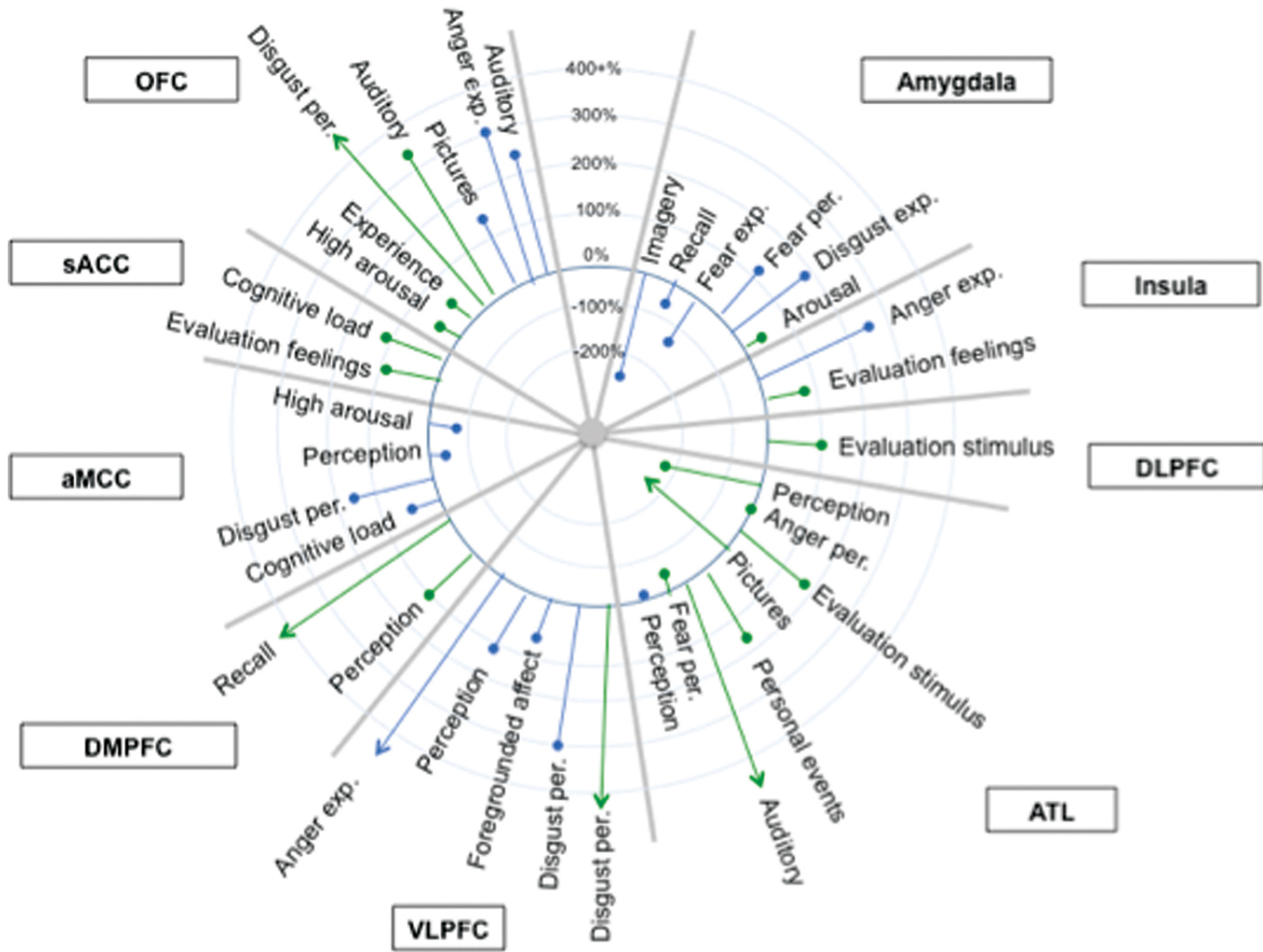
...emotions emerge when people make meaning out of sensory input from the body and from the world using knowledge of prior experiences. Emotions are “situated conceptualizations” (cf. Barsalou 2003) because the emerging meaning is tailored to the immediate environment and prepares the person to respond to sensory input in a way that is tailored to the situation



(Lindquist, Wager, Kober, Bliss-Moreau, & Barrett, 2012)
(<http://dx.doi.org/10.1017/S0140525X11000446>)

Figure 4. The Neural Reference Space for Discrete Emotion. The neural reference space (phrase coined by Edelman [1989]) is the set of brain regions consistently activated across all studies assessing the experience or perception of anger, disgust, fear, happiness and sadness (i.e. the superordinate category emotion). Brain regions in yellow exceeded the height threshold ($p < .05$) and regions in orange exceeded the most stringent extent-based threshold ($p < .001$). Regions in pink and magenta correspond to lesser extent-based thresholds and are not discussed in this article. Cortex is grey, the brainstem and nucleus accumbens are green, the amygdala is blue and the cerebellum is purple. A color version of this image can be viewed in the online version of this target article at <http://www.journals.cambridge.org/bbs> (<http://www.journals.cambridge.org/bbs>).

(Lindquist, Wager, Kober, Bliss-Moreau, & Barrett, 2012)
(<http://dx.doi.org/10.1017/S0140525X11000446>)



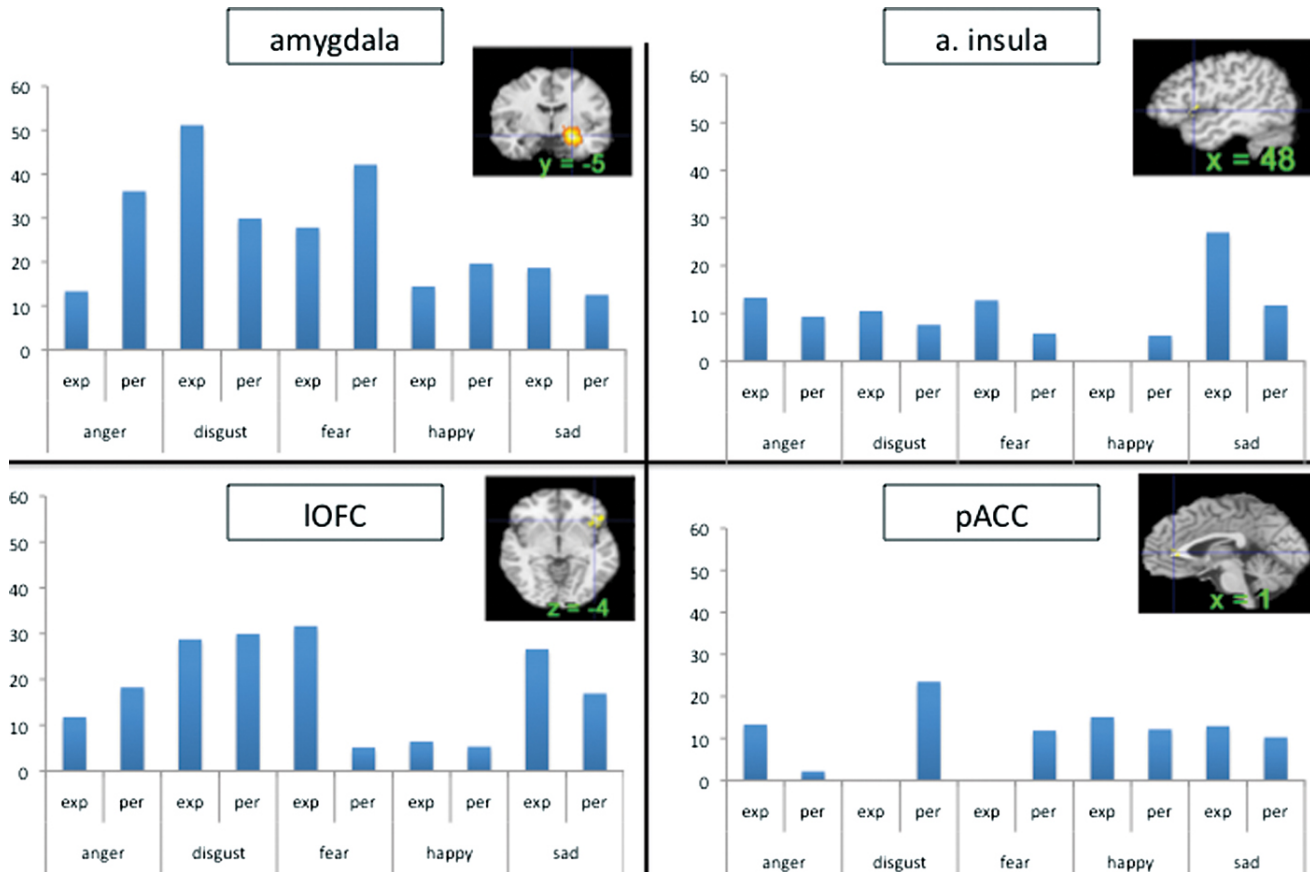
(Lindquist, Wager, Kober, Bliss-Moreau, & Barrett, 2012)

(<http://dx.doi.org/10.1017/S0140525X11000446>)

Figure 5. Logistic Regression Findings. Selected results from the logistic regressions are presented (for additional findings, see Table S6 in supplementary materials). Circles with positive values represent a 100% increase in the odds that a variable predicted an increase in activity in that brain area. Circles with negative values represent a 100% increase in the odds that a variable predicted there would not be an increase in activity in that brain area.

Legend: Blue lines: left hemisphere; Green lines: right hemisphere. Arrowheads: % change in odds is greater than values represented in this figure. Abbreviations: OFC: orbitofrontal cortex; DLPFC: dorsolateral prefrontal cortex; ATL: anterior temporal lobe; VLPFC: ventrolateral prefrontal cortex; DMPFC: dorsomedial prefrontal cortex; aMCC: anterior mid-cingulate cortex; sAAC: subgenual ACC. A color version of this image can be viewed in the online version of this target article at <http://www.journals.cambridge.org/bbs> (<http://www.journals.cambridge.org/bbs>).

(Lindquist, Wager, Kober, Bliss-Moreau, & Barrett, 2012)
(<http://dx.doi.org/10.1017/S0140525X11000446>)



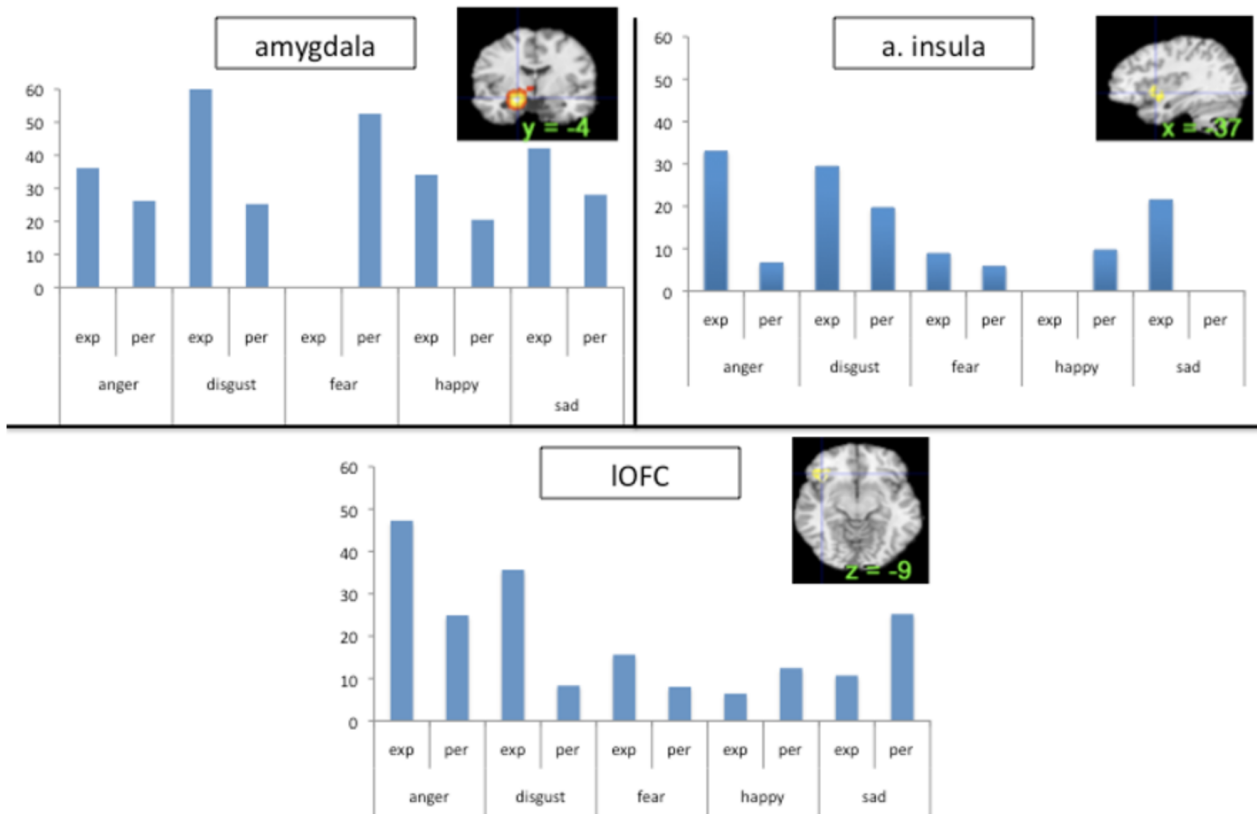
(Lindquist, Wager, Kober, Bliss-Moreau, & Barrett, 2012)

(<http://dx.doi.org/10.1017/S0140525X11000446>)

Figure 6. Proportion of Study Contrasts with Increased Activation in Four Key Brain Areas. The y-axes plot the proportion of study contrasts in our database that had increased activation within 10mm of that brain area. The x-axes denote the contrast type separated by experience (exp) and perception (per). All brain regions depicted are in the right hemisphere. See Figures S2 and S3 in supplementary materials, available at
<http://www.journals.cambridge.org/bbs2012008>
(<http://www.journals.cambridge.org/bbs2012008>), for additional regions. A color version of this image can be viewed in the online version of this target article at
<http://www.journals.cambridge.org/bbs>
(<http://www.journals.cambridge.org/bbs>).

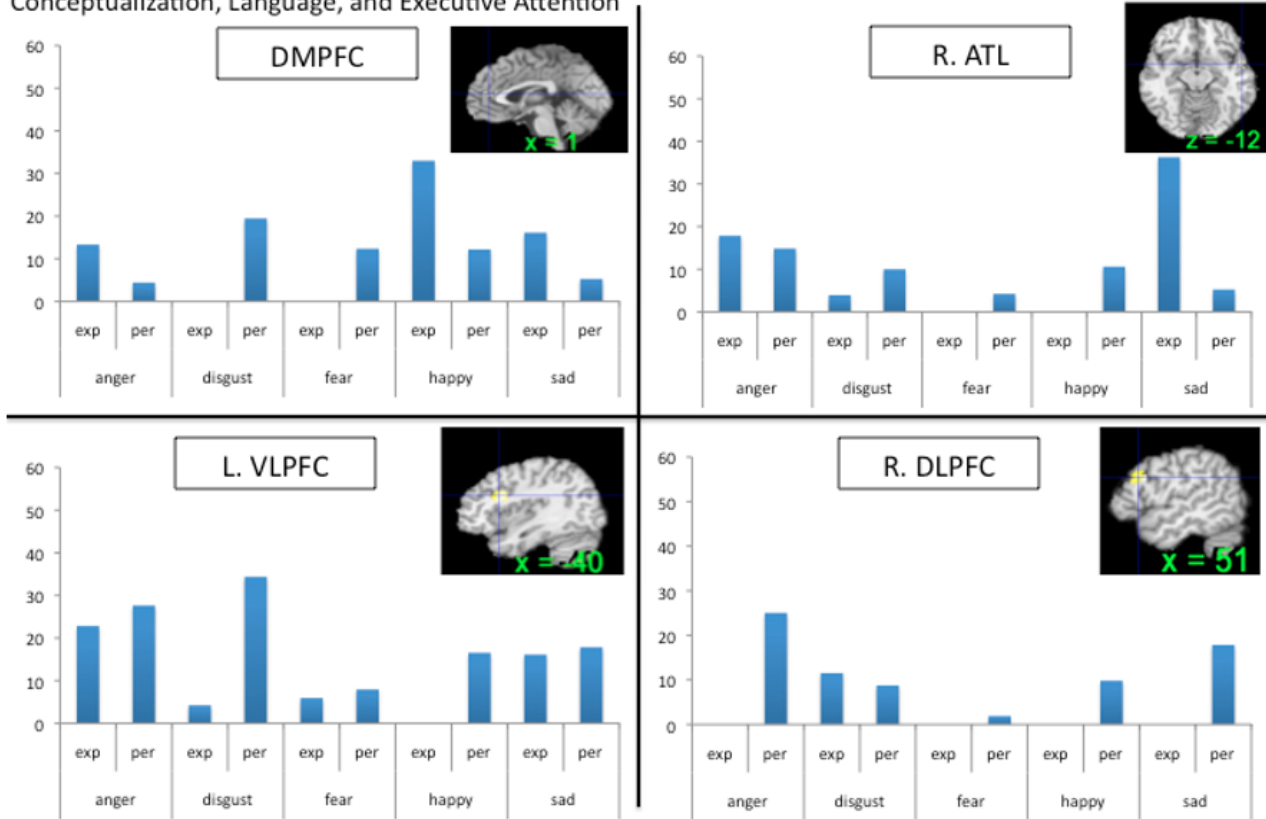
(Lindquist, Wager, Kober, Bliss-Moreau, & Barrett, 2012)
(<http://dx.doi.org/10.1017/S0140525X11000446>)

Figure S2. Proportion of Study Contrasts with Increased Activation in Key Brain Areas



(Lindquist, Wager, Kober, Bliss-Moreau, & Barrett, 2012)
<http://dx.doi.org/10.1017/S0140525X11000446>)

Figure S3. Proportion of Study Contrasts with Increased Activation in Brain Regions associated with Conceptualization, Language, and Executive Attention



The y-axes plot the proportion of study contrasts in our database that had increased activation within 10mm of that brain area.

(Lindquist, Wager, Kober, Bliss-Moreau, & Barrett, 2012)
<http://dx.doi.org/10.1017/S0140525X11000446>)

Amygdala as a ‘hub’ for fear

Our meta-analytic findings were inconsistent with a locationist hypothesis of amygdala function but were more consistent with the psychological constructionist hypothesis. Our density analyses revealed that, as compared to other brain regions, voxels within both amygdalae had more consistent increases in activation during instances of fear perception than during the perception of any other emotion category (Table 1). These voxels were not functionally specific for instances of perceiving fear, however.

Anterior insula as ‘hub’ for disgust

Our meta-analytic findings were inconsistent with the locationist account that the anterior insula is the brain seat of disgust but were more consistent with the psychological constructionist account that insula activity is correlated with interoception and the awareness of affective feelings.

Orbitofrontal cortex (OFC) as ‘hub’ for anger

Our meta-analytic findings were inconsistent with the locationist hypothesis that the OFC is the brain seat of anger. As compared to voxels within other brain regions, voxels within the OFC did not have more consistent increases during instances of anger experience or perception than during any other emotion category. Rather, as compared to voxels within other brain regions, voxels within the left IOFC had more consistent increases in activation during instances of disgust experience than during the experience of other emotion categories.

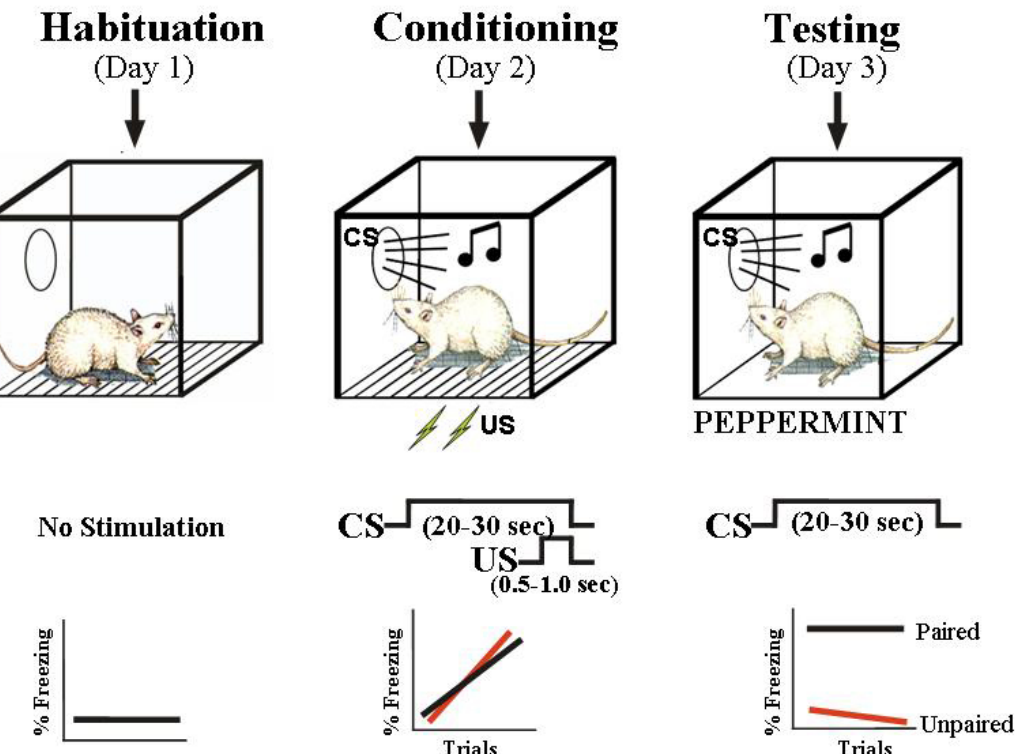
Anterior cingulate cortex as ‘hub’ for sadness

Our meta-analytic evidence is inconsistent with the locationist account that the ACC is the brain basis of sadness, but more consistent with a psychological constructionist hypothesis of ACC function. As compared to voxels within other brain regions, voxels within the sACC, pACC and aMCC did not have more consistent increases when participants were experiencing or perceiving instances of sadness than during any other emotion category (Fig. 6).

Fear

Animal models

Pavlovian Threat Conditioning Paradigm



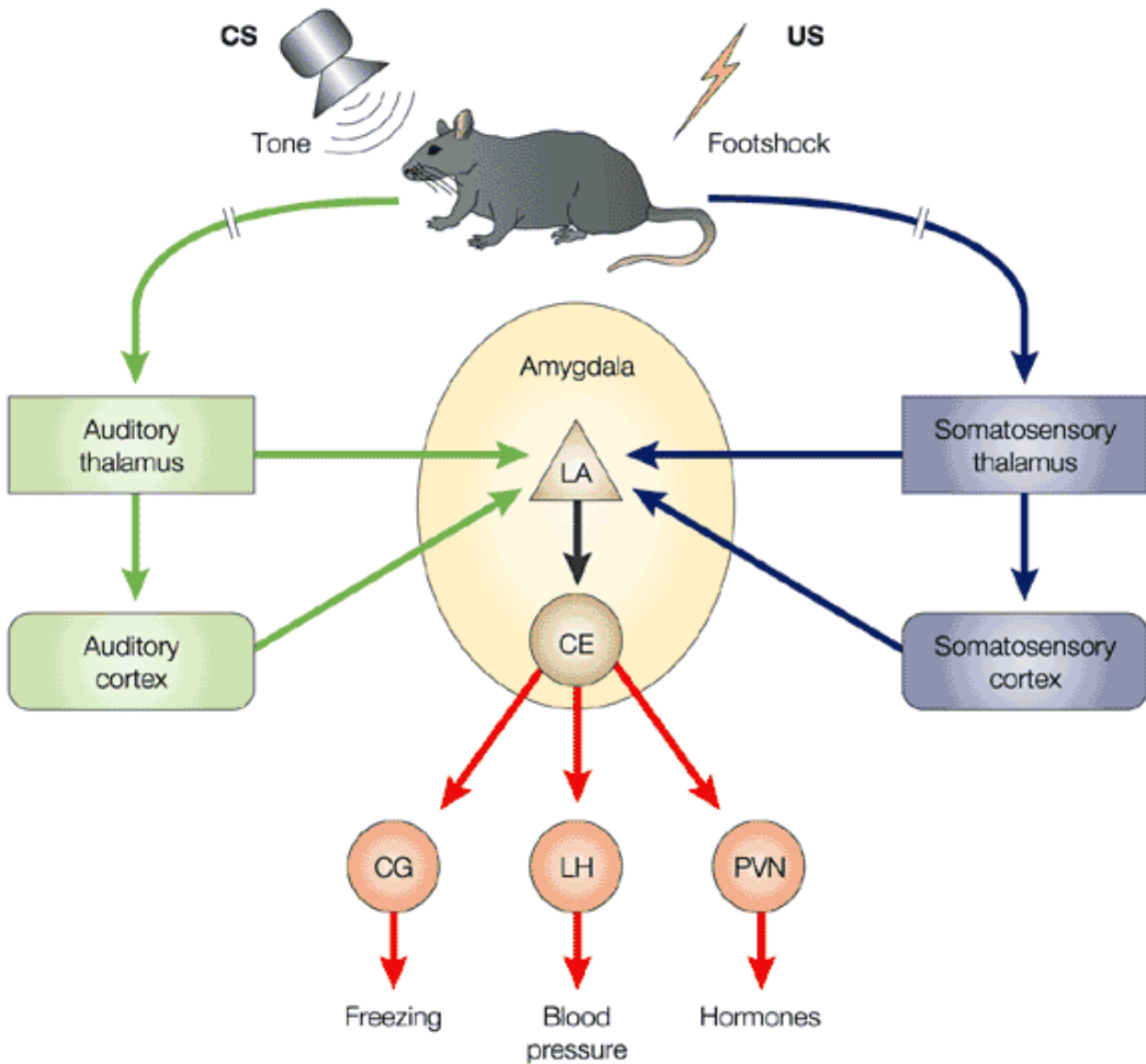
http://www.cns.nyu.edu/labs/ledouxlab/images/image_research/fear_conditioning.jpg
[\(http://www.cns.nyu.edu/labs/ledouxlab/images/image_research/fear_conditioning.jpg\)](http://www.cns.nyu.edu/labs/ledouxlab/images/image_research/fear_conditioning.jpg)

<https://youtu.be/ZlZekx1P1g4> (<https://youtu.be/ZlZekx1P1g4>)

Measures in Animal Model	DSM-III: Generalized Anxiety
Heart rate increase	Heart pounding
Salivation decrease	Dry mouth
Stomach ulcers	Upset stomach
Respiration change	Respiration increase
Scanning & vigilance	Scanning & vigilance
Startle response increase	Jumpiness, easy startle
Urination	Frequent urination
Defecation	Diarrhea
Grooming	Fidgeting
Freezing	Apprehensive expectation

Adapted from (Davis, 1992) ([http://dx.doi.org/10.1016/0165-6147\(92\)90014-W](http://dx.doi.org/10.1016/0165-6147(92)90014-W))

Amygdala circuits



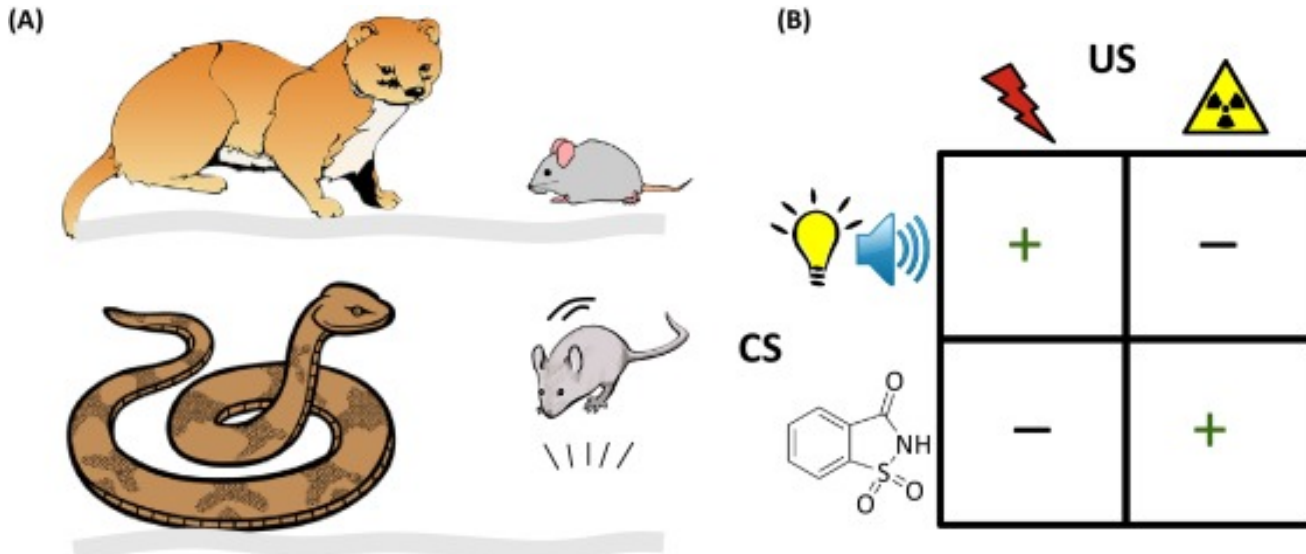
Nature Reviews | Neuroscience

(Medina, Repa, Mauk, & LeDoux, 2002)

(<http://dx.doi.org/10.1038/nrn728>)

- Direct (fast) pathways via thalamus
- Indirect (slower) pathways via cortex
- Input and output (behavior, physiology) specificity

Specificity of learning stimulus/response mappings



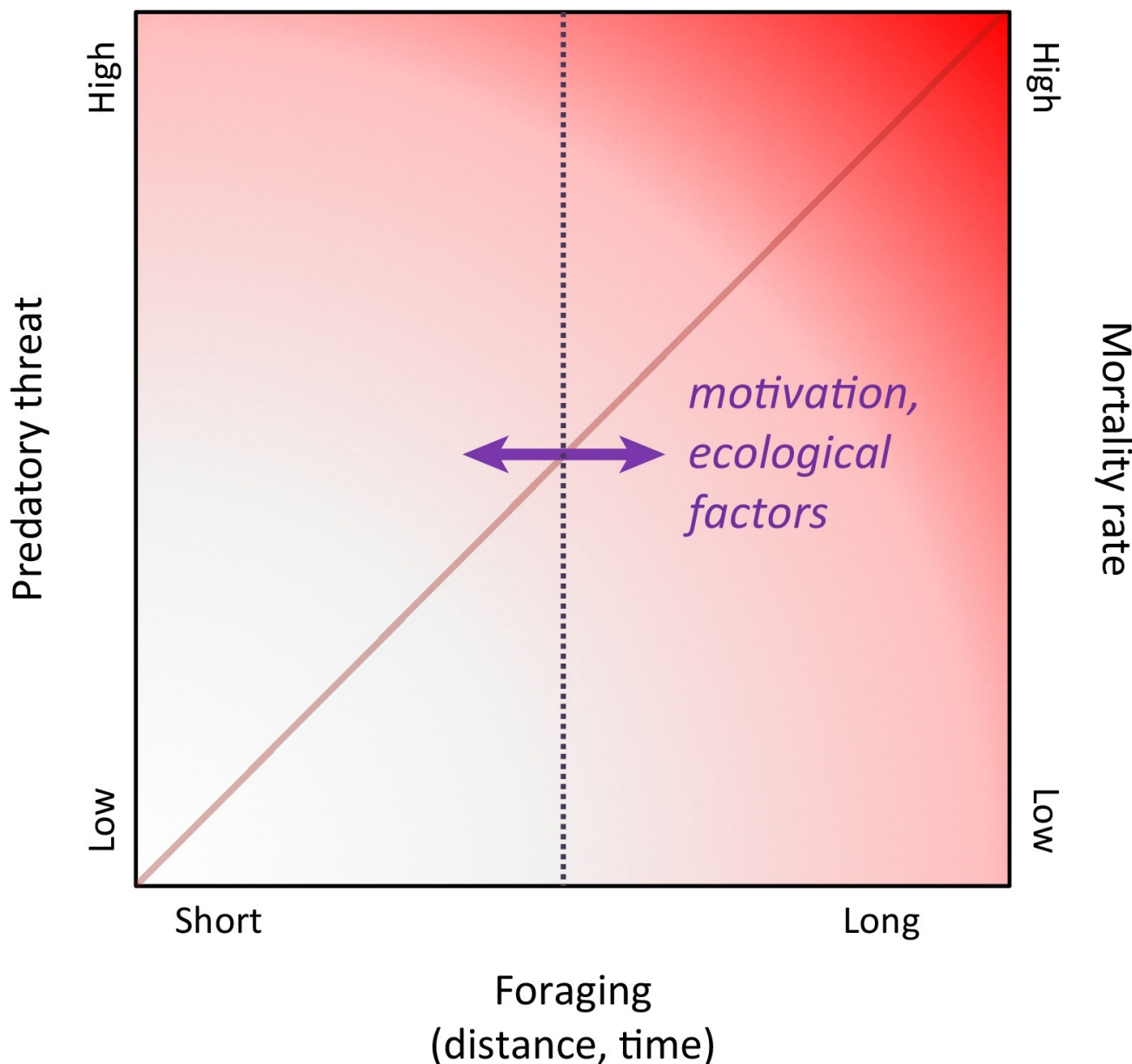
Trends in Neurosciences

(Pellman & Kim, 2016) (<https://doi.org/10.1016/j.tins.2016.04.001>)

Figure 1. Evolutionary Influences on Innate and Learned Fear. (A) Predatory history shapes prey's innate fear responses as illustrated by *Peromyscus maniculatus austerus* deer mouse's freezing to weasels and *Peromyscus maniculatus gambeli* deer mouse's jump (Jan Gillbank, 'Drawing of a grey mouse' October 27, 2012 via Wikimedia, Creative Commons Attribution 3.0 License) to gopher snakes [2]. *P. m. austerus* deer mice live in the coniferous forests of western Washington State and *P. m. gambeli* deer mice dwell in the arid grassland of eastern Washington State. (B) Ecological history predisposes fear learning. A classic study by John Garcia [3] found that rats easily acquired conditioned fear to bright/noisy conditioned stimulus (CS) paired to footshock unconditioned stimulus (US) and conditioned taste aversion to saccharin taste CS paired to X-rays (or LiCl) US. However, rats showed lack of conditioning to bright/noisy-X-ray (or LiCl) and saccharin-footshock pairings.

(Pellman & Kim, 2016) (<https://doi.org/10.1016/j.tins.2016.04.001>)

- Specific stimulus/response, $S \rightarrow R$, patterns
- Visual OR Auditory \rightarrow pain
- Taste \rightarrow nausea



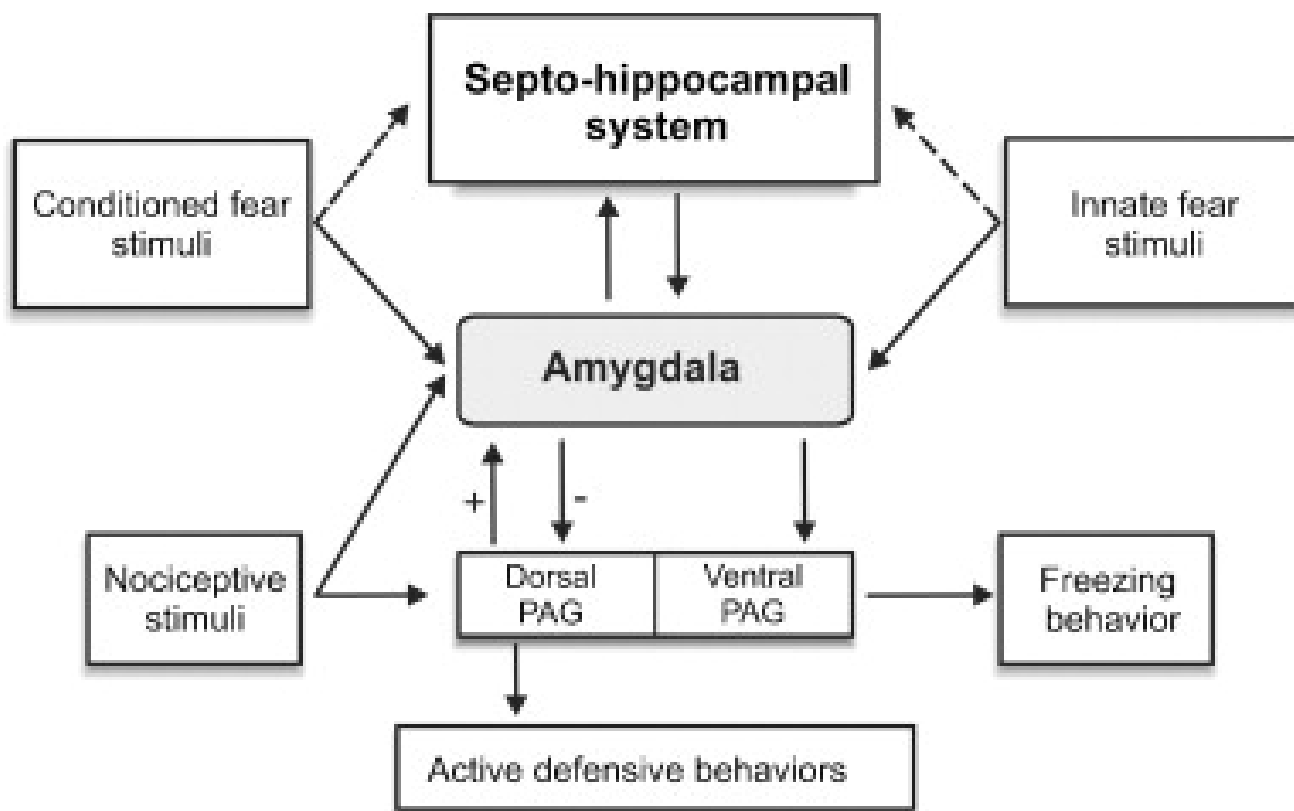
Trends in Neurosciences

(Pellman & Kim, 2016) (<https://doi.org/10.1016/j.tins.2016.04.001>)

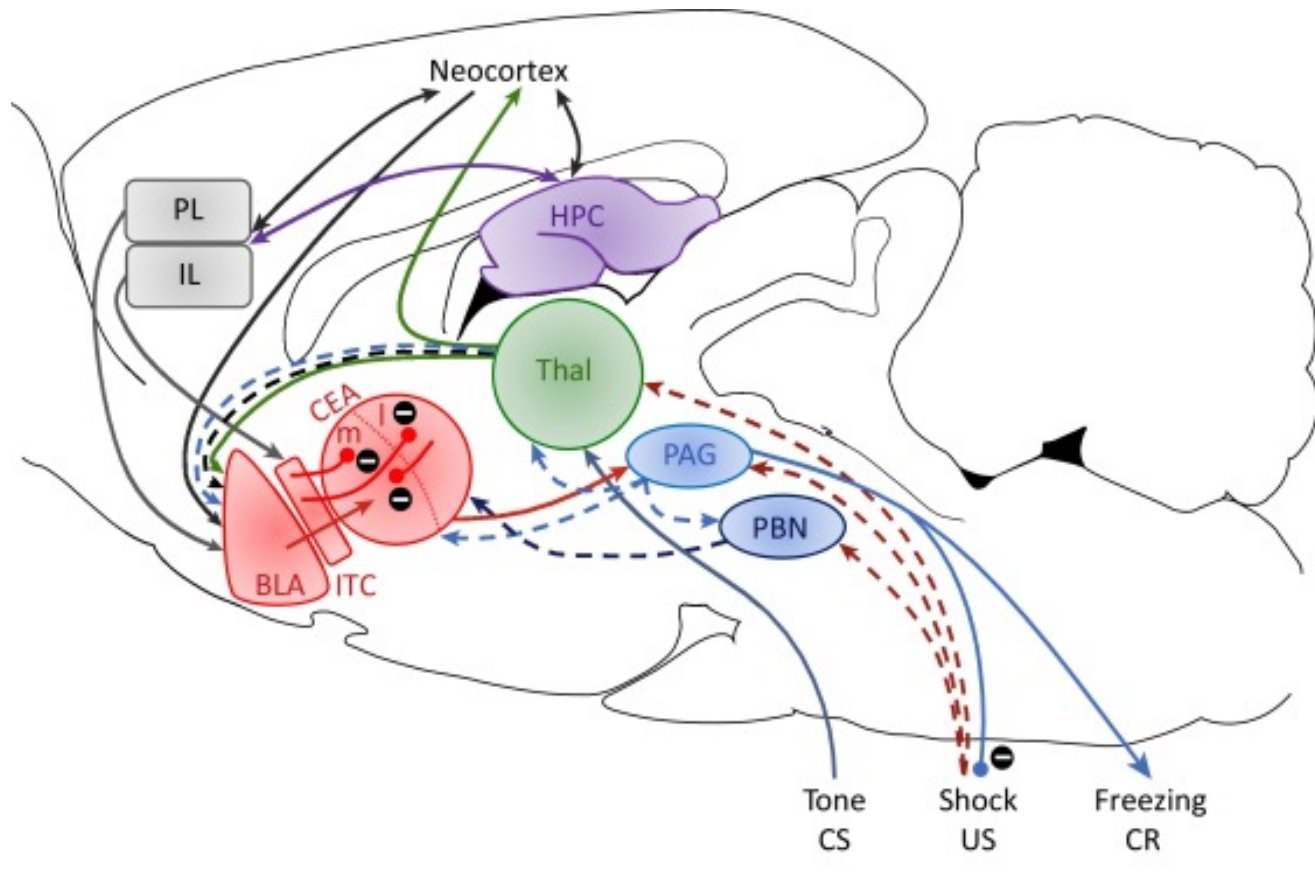
Figure 4. Foraging and Risk of Predation. Foraging distance and time away from the safety of a nest are positively correlated with the risk of meeting predators, which can result in injury or death. Motivational factors, such as hunger, reproductive and parental state, and ecological factors, such as food availability and predator density, influence foraging behavior (represented by a horizontal arrow) and thus predation risk. Fear elicits immediate species-specific defense reactions upon meeting a predator and exerts enduring influences on foraging strategy.

(Pellman & Kim, 2016) (<https://doi.org/10.1016/j.tins.2016.04.001>)

Circuitry



(Brandão, Zanoveli, Ruiz-Martinez, Oliveira, & Landeira-Fernandez, 2008) (<http://dx.doi.org/10.1016/j.bbr.2007.10.018>)



Trends in Neurosciences

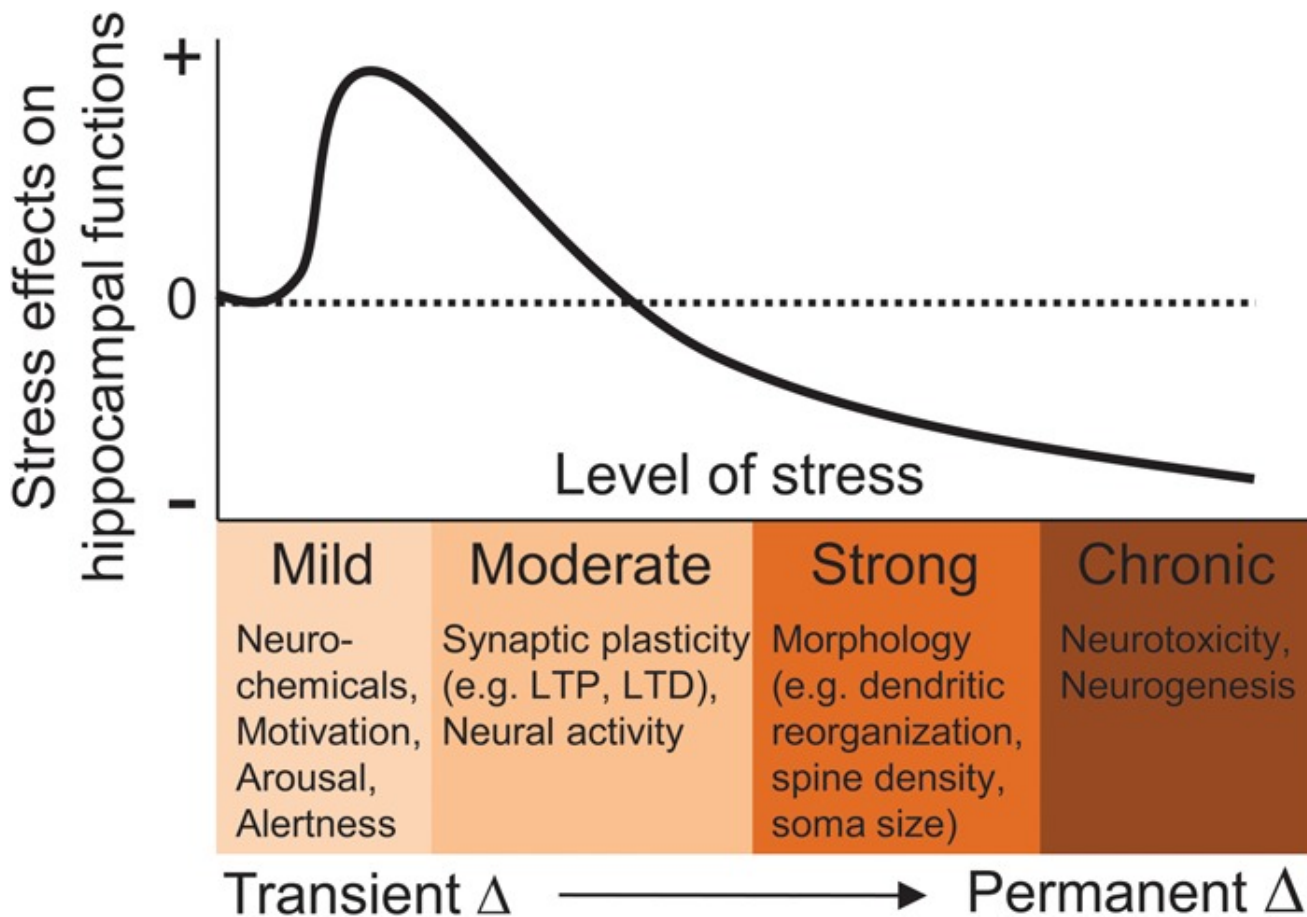
(Pellman & Kim, 2016) (<https://doi.org/10.1016/j.tins.2016.04.001>)

- BLA, basolateral complex of the amygdala
- CEA, central nucleus of the amygdala
- ITC, intercalated cells of the amygdala
- PL, prelimbic cortex
- IL, infralimbic cortex
- HPC, hippocampus
- Thal, thalamus
- PAG, periaqueductal gray
- PBN, parabrachial nucleus

Stress types

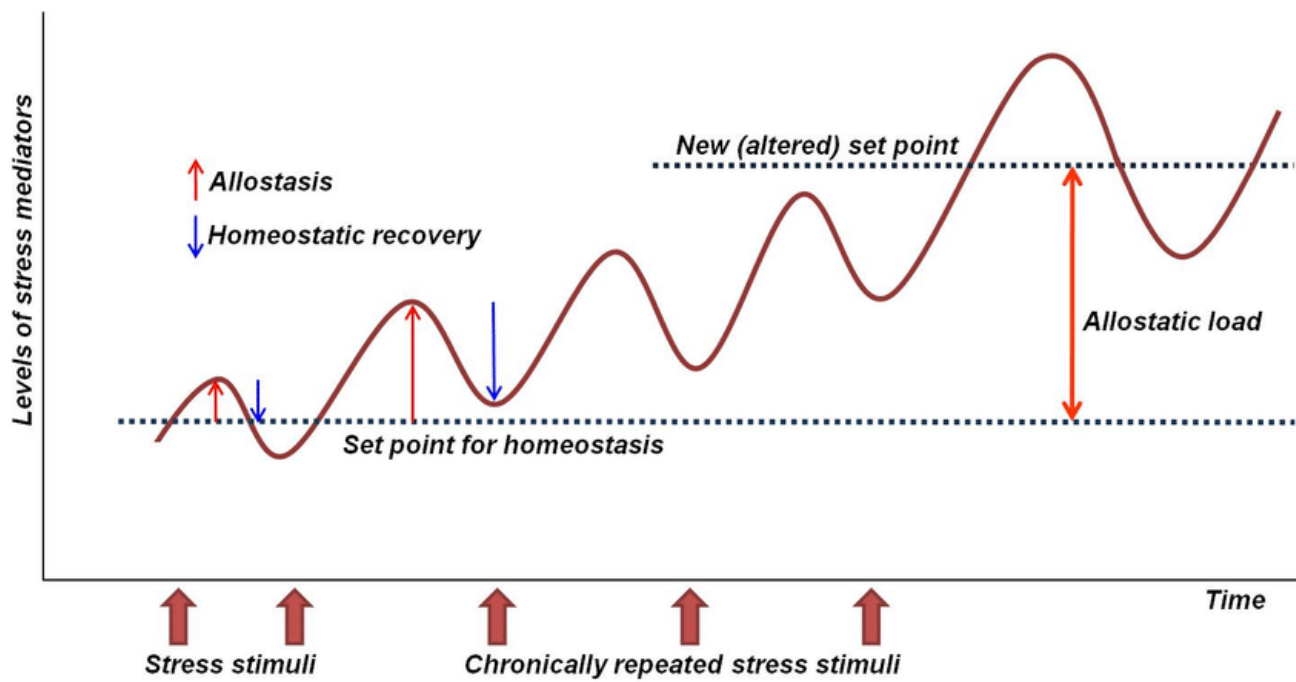
- **Acute** stress
 - Short duration
- Brain detects threat

- Mobilizes physiological, behavioral responses
 - HPA (Cortisol), SAM (NE/Epi) axes
- vs. **Chronic** or stress
 - Long duration, persistent



(Kim, Pellman, & Kim, 2015) (<http://dx.doi.org/10.1101/lm.037291.114>)

- Homeostasis vs. allostasis

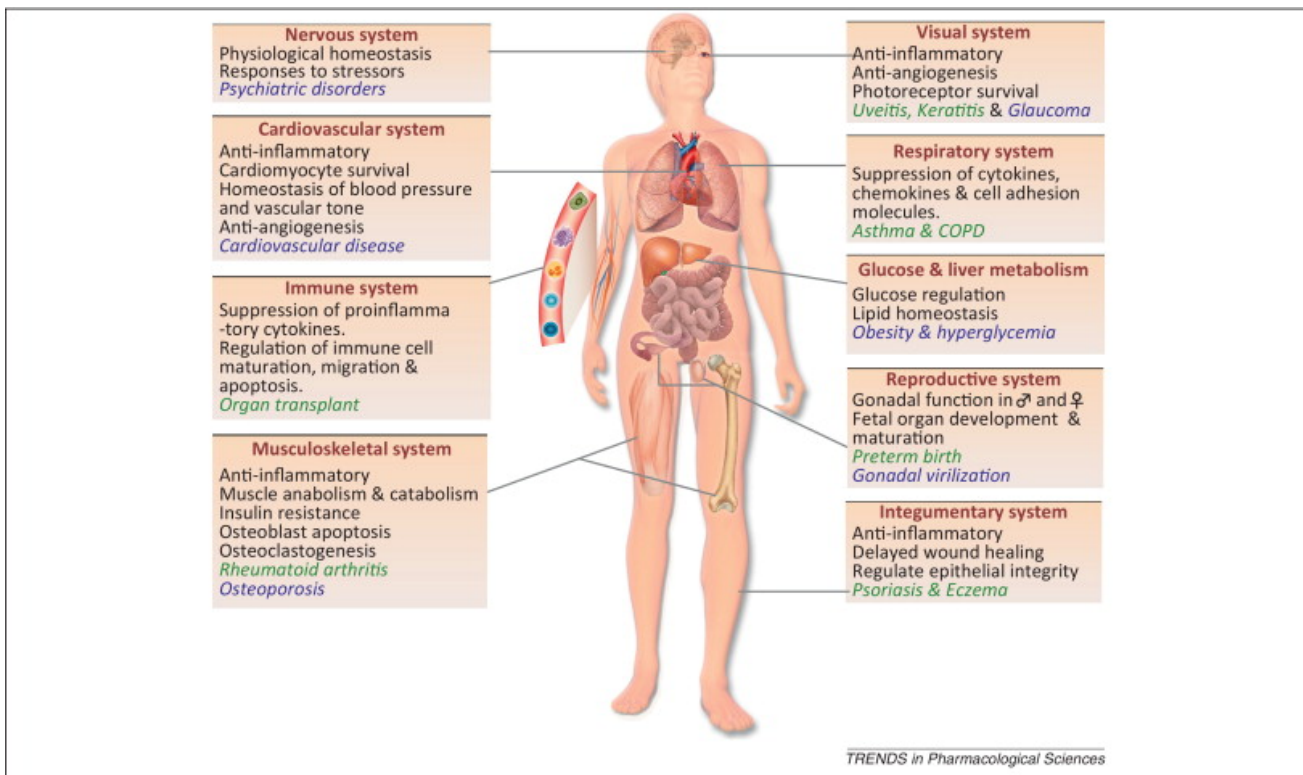


(Lee, Kim, & Choi, 2015)

(<http://dx.doi.org/10.5483/bmbrep.2015.48.4.275>)

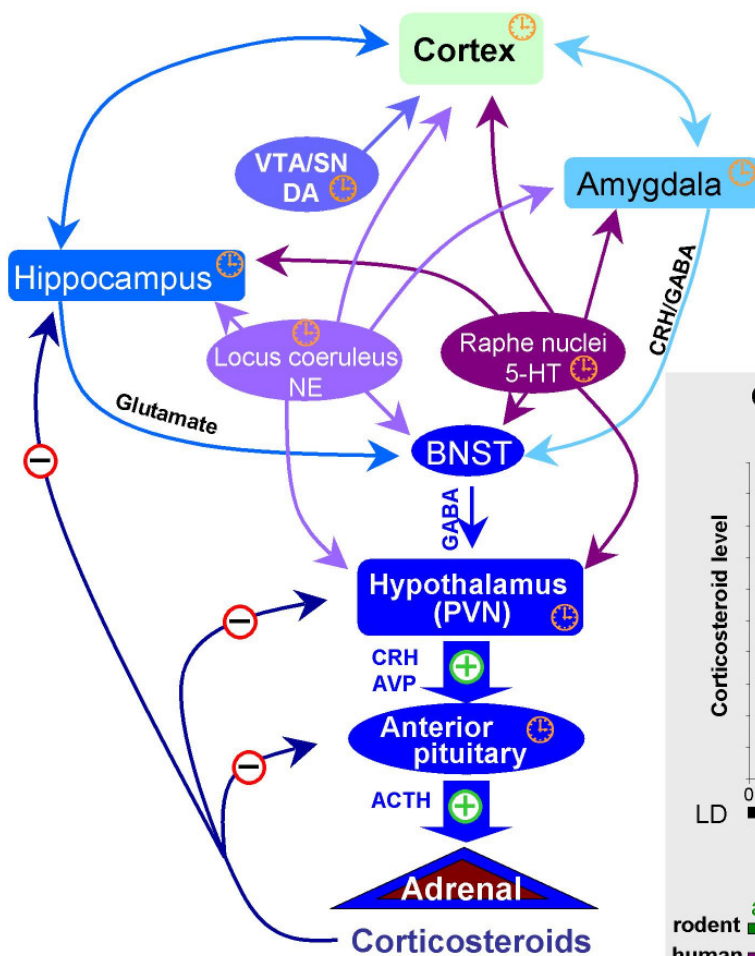
Glucocorticoids

- Released by
 - Adrenal cortex
 - Other areas in small amounts
- Cortisol (hydrocortisone)
 - Increases blood glucose levels
 - Aids in fat, protein, carbohydrate metabolism
 - Suppresses immune system
 - Reduces inflammation
- Receptors in body and brain



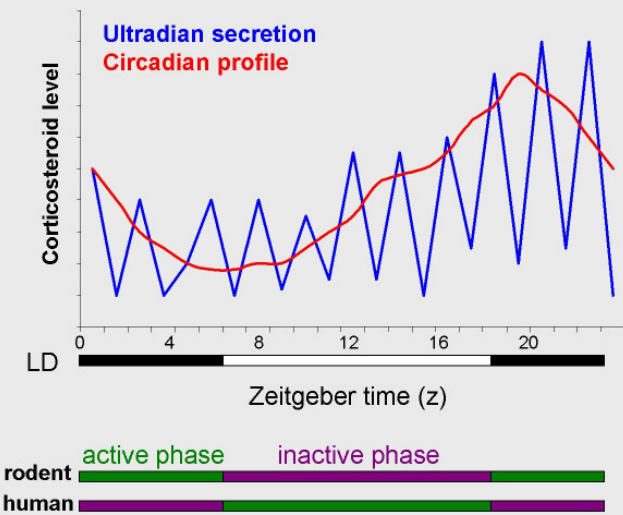
(Kadmiel & Cidlowski, 2013) (10.1016/j.tips.2013.07.003)

- Multiple feedback loops
- Diurnal pattern



- rapid behavioral responses**
- memory consolidation and retrieval
 - fear and anxiety
 - aggression
 - locomotion
 - vigilance and gating
 - reward

Circadian and ultradian rhythms of corticosteroid secretion

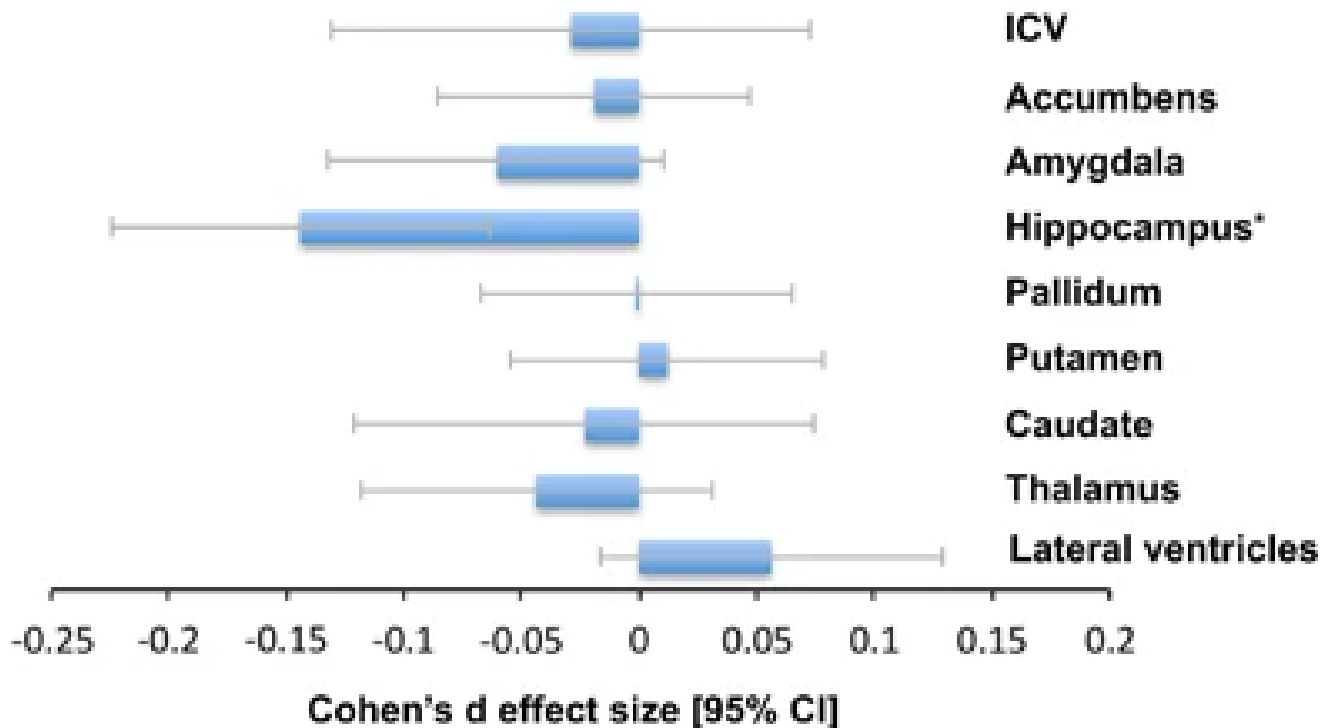


<http://www.molecularbrain.com/content/figures/1756-6606-3-2-1-l.jpg>
[\(http://www.molecularbrain.com/content/figures/1756-6606-3-2-1-l.jpg\)](http://www.molecularbrain.com/content/figures/1756-6606-3-2-1-l.jpg)

Impacts of chronic stress

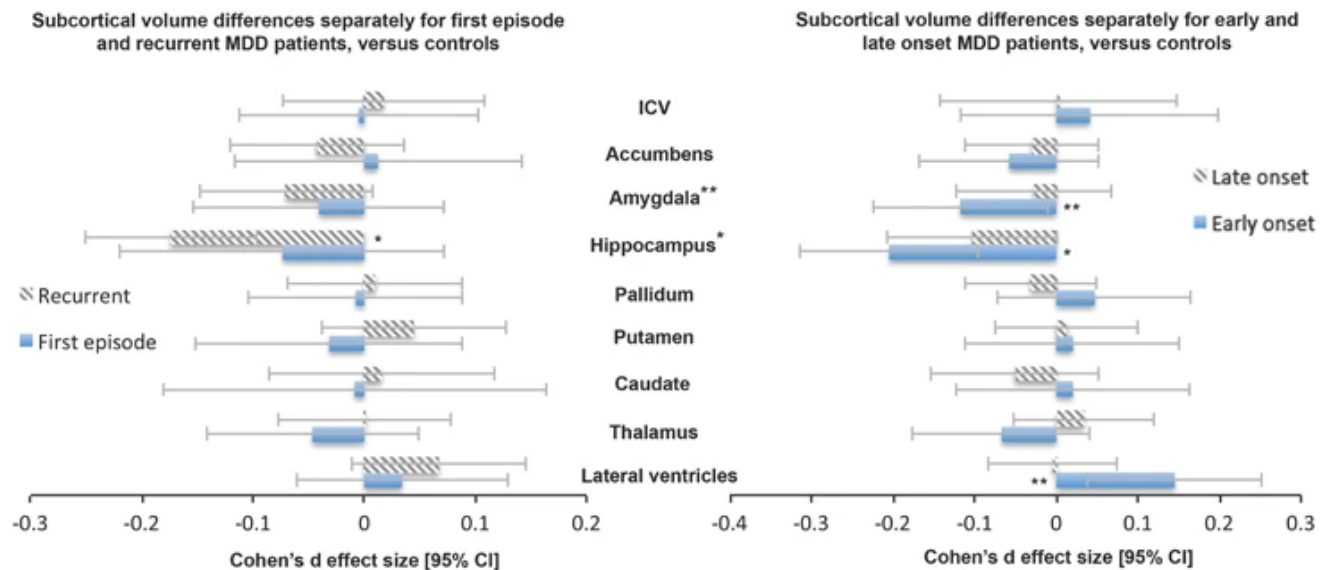
- Major depressive disorder (MDD) & Post-traumatic Stress Disorder (PTSD)
 - Hippocampus and PFC volume reductions
 - Synapse loss
 - Reduced dendritic density

Subcortical volume differences MDD patients and controls



(Schmaal et al., 2016) (<http://dx.doi.org/10.1038/mp.2015.69>)

Cohen's d-effect sizes 95% CI and for differences in subcortical brain volumes between major depressive disorder (MDD) patients and healthy control subjects. Effect sizes were corrected for age, sex and intracranial volume (ICV). The effect size for ICV was corrected for age and sex. P<0.05 corrected. CI, confidence interval.

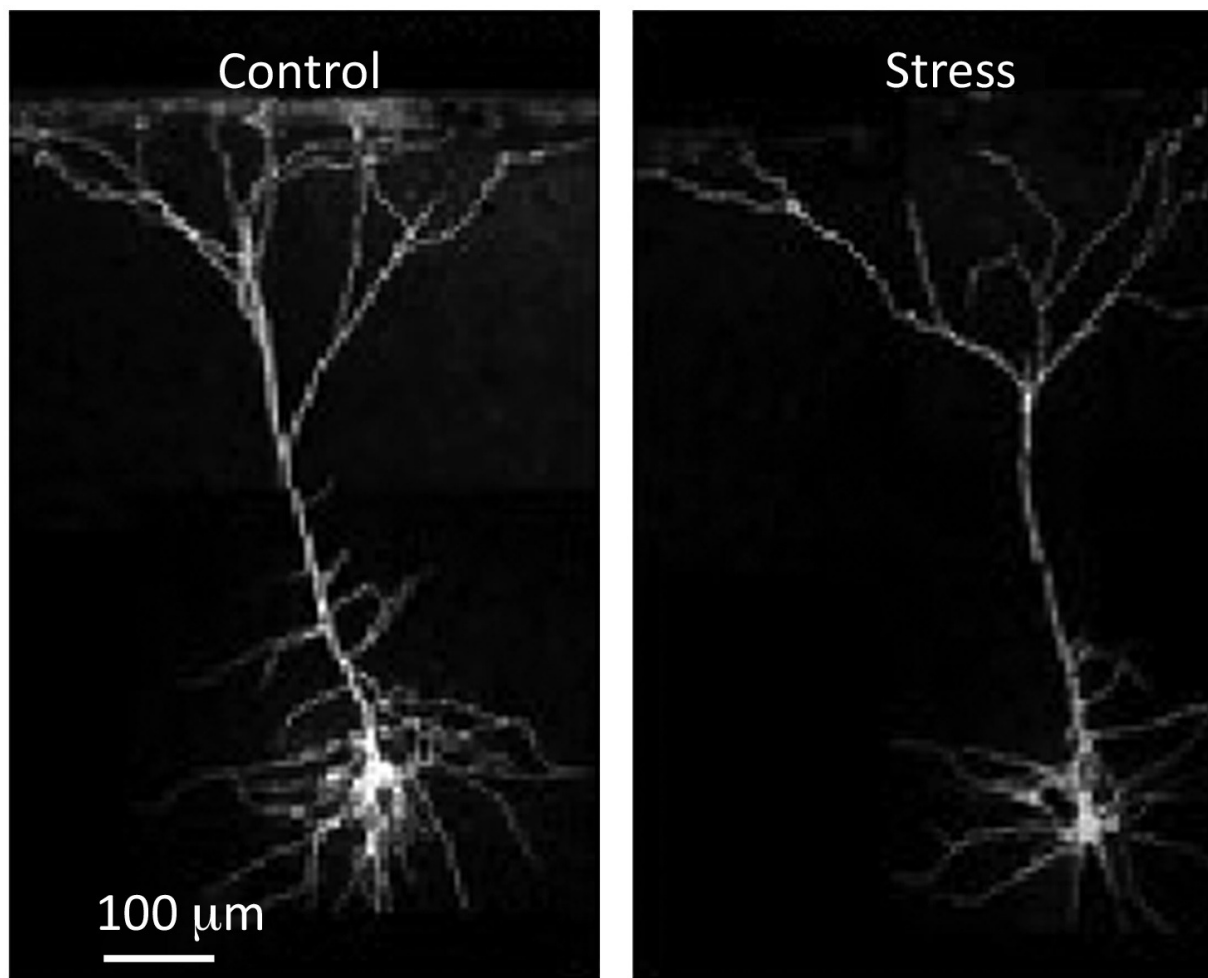


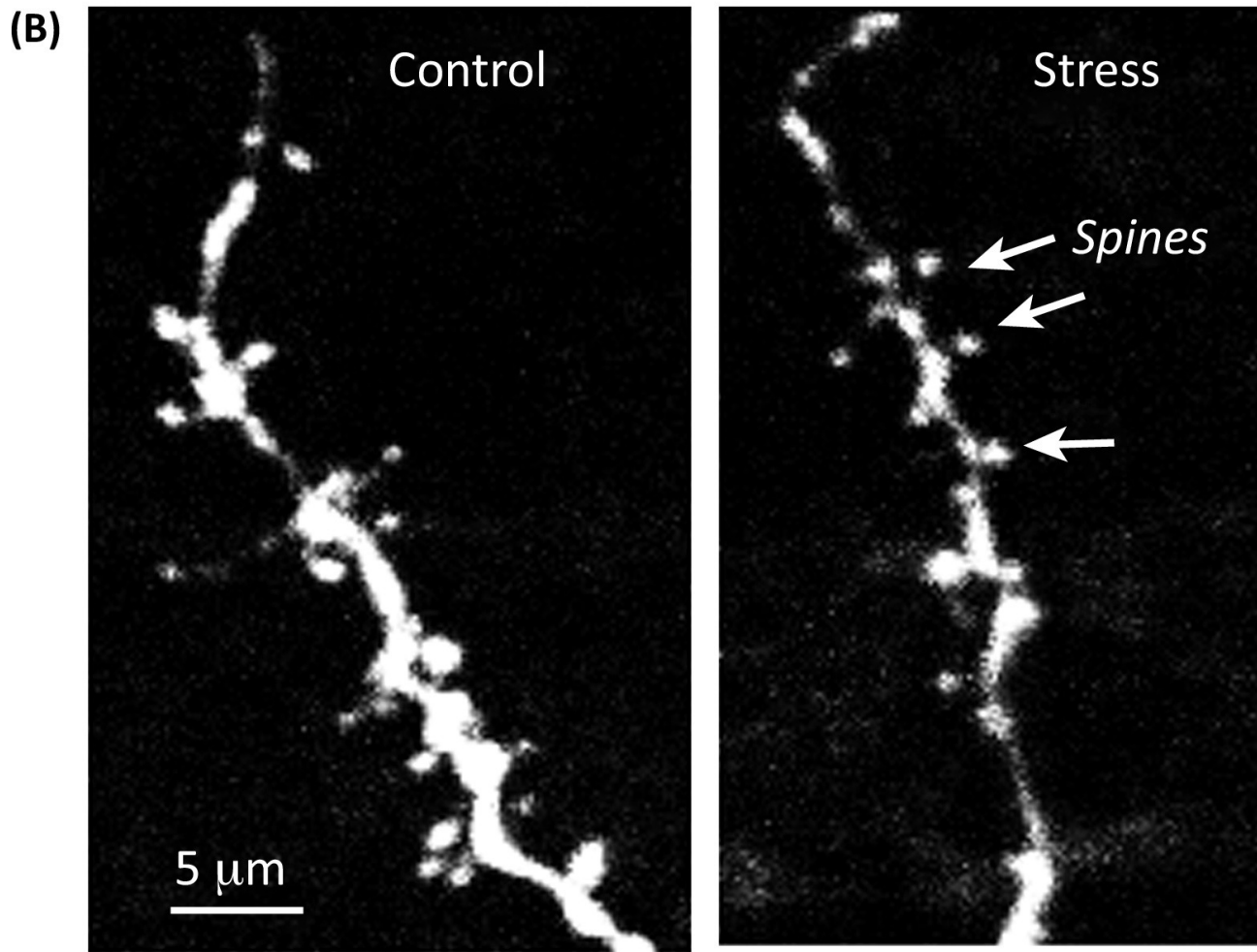
(Schmaal et al., 2016) (<http://dx.doi.org/10.1038/mp.2015.69>)

(a) Cohen's d -effect sizes 95% CI for differences in subcortical brain volumes between recurrent major depressive disorder (MDD) patients and healthy control subjects (striped pattern) and between first episode MDD patients and healthy controls (no pattern). (b) Cohen's d -effect sizes 95% CI for differences in subcortical brain volumes between early onset (≤ 21) MDD patients and healthy control subjects (no pattern) and between later onset (> 21) MDD patients and healthy controls (striped pattern). Effect sizes were corrected for age, sex and intracranial volume (ICV). $P < 0.05$ corrected, $P < 0.05$. CI, confidence interval.

Impacts of acute stress

(A)







0 stress

1x stress

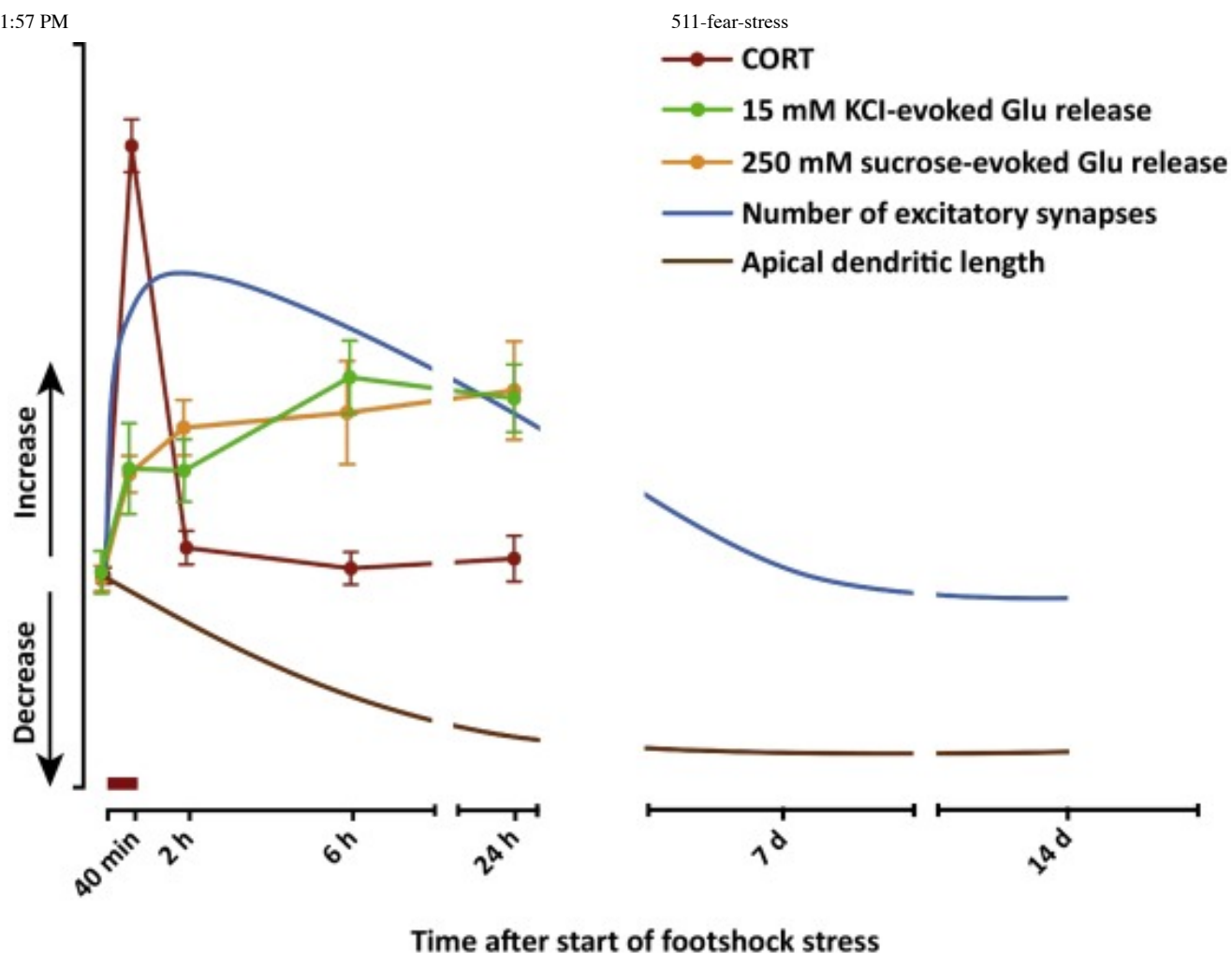
3x stress

Trends in Neurosciences

(Musazzi, Tornese, Sala, & Popoli, 2017)

<https://doi.org/10.1016/j.tins.2017.07.002>

“Figure 1. Neuroarchitectural Changes Induced by Repeated or Acute Stress in Rodents. (A) Repeated restraint stress (7 days) induces a reduction in the number and length of apical dendrites of pyramidal neurons (layer V) in the medial prefrontal cortex (PFC) of rats. (B) Magnified segment of dendrite from the same stressed rats, showing that repeated stress significantly decreases the number of spine synapses in medial PFC. (C) Reconstructions of representative infralimbic pyramidal neurons in mice exposed to zero (0), one (1×), or three (3×) unpredictable sessions of 10 min of forced swim stress. Apical dendritic branch length was significantly reduced after one or three stress episodes relative to controls. Adapted, with permission, from [24] (B) and [23] (C).”



Trends in Neurosciences

(Musazzi, Tornese, Sala, & Popoli, 2017)
<https://doi.org/10.1016/j.tins.2017.07.002>

Figure 3. Graphic Summary of Short- and Long-Term Functional and Neuroarchitectural Effects in Prefrontal Cortex (PFC) Synapses after Acute Footshock (FS) Stress [44]. The fast and transient increase in corticosterone (CORT) release induced by acute (40 min) FS stress was accompanied by the rapid increase in both depolarization-evoked and hypertonic sucrose-evoked (readily releasable pool) glutamate release in PFC, and the number of small excitatory synapses. The enhancement of glutamate release was sustained for up to 24 h, as well as the increased number of excitatory synapses, which normalized between 24 h and 7 days after FS. Before 24 h had elapsed from the start of FS stress, retraction of apical dendrites began and was sustained for up to 14 days. The timing of actual FS stress (40 min) is indicated by the red marker. Number of excitatory synapses and apical dendrite length are indicative and not in scale with other readouts. CORT and glutamate release data adapted from [44].

Changes in neural architecture

- Hippocampus (rich in CORT receptors)
- Prefrontal cortex

Neurochemical factors

- Cortisol enhances glutamate release
- Corticosteroid antagonists block this
- Ketamine (NMDA receptor antagonist) may act via similar mechanisms

ROBERT M. SAPOLSKY

Author of *A Primate's Memoir*

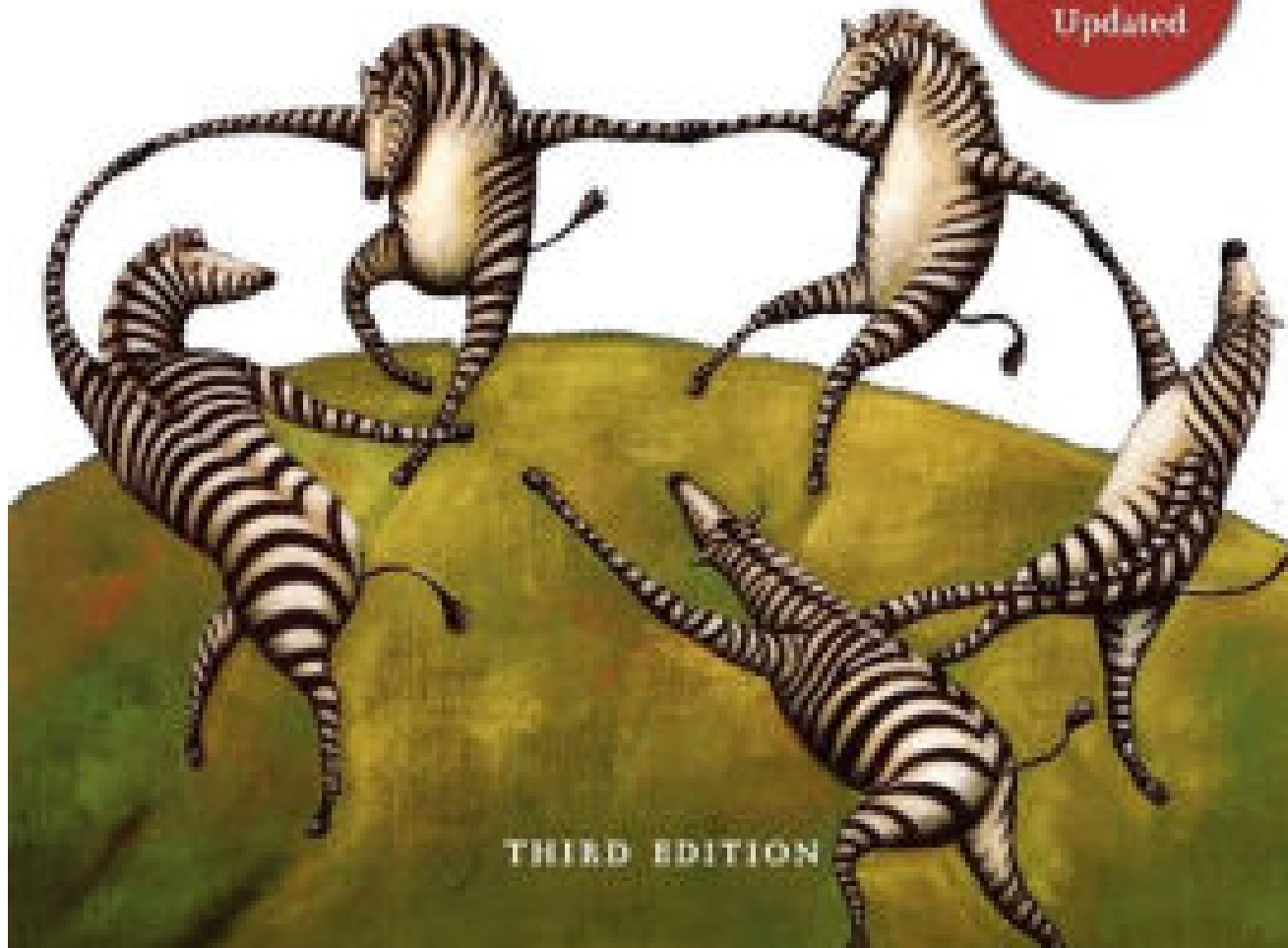
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References

- Brandão, M. L., Zanoveli, J. M., Ruiz-Martinez, R. C., Oliveira, L. C., & Landeira-Fernandez, J. (2008). Different patterns of freezing behavior organized in the periaqueductal gray of rats: Association with different types of anxiety. *Behavioural Brain Research*, 188(1), 1–13. [\(https://doi.org/10.1016/j.bbr.2007.10.018\)](https://doi.org/10.1016/j.bbr.2007.10.018)
- Davis, M. (1992). The role of the amygdala in fear-potentiated startle: Implications for animal models of anxiety. *Trends in Pharmacological Sciences*, 13, 35–41. [\(https://doi.org/10.1016/0165-6147\(92\)90014-W\)](https://doi.org/10.1016/0165-6147(92)90014-W)
- Kadmiel, M., & Cidlowski, J. A. (2013). Glucocorticoid receptor signaling in health and disease. *Trends in Pharmacological Sciences*, 34(9), 518–530. [\(https://doi.org/10.1016/j.tips.2013.07.003\)](https://doi.org/10.1016/j.tips.2013.07.003)
- Kim, E. J., Pellman, B., & Kim, J. J. (2015). Stress effects on the hippocampus: A critical review. *Learning & Memory*, 22(9), 411–416. [\(https://doi.org/10.1101/lm.037291.114\)](https://doi.org/10.1101/lm.037291.114)
- Lee, D. Y., Kim, E., & Choi, M. H. (2015). Technical and clinical aspects of cortisol as a biochemical marker of chronic stress. *BMB Reports*, 48(4), 209–216. [\(https://doi.org/10.5483/bmbrep.2015.48.4.275\)](https://doi.org/10.5483/bmbrep.2015.48.4.275)
- Lindquist, K. A., Wager, T. D., Kober, H., Bliss-Moreau, E., & Barrett, L. F. (2012). The brain basis of emotion: A meta-analytic review. *Behav. Brain Sci.*, 35(3), 121–143. [\(https://doi.org/10.1017/S0140525X11000446\)](https://doi.org/10.1017/S0140525X11000446)

- Medina, J. F., Repa, J. C., Mauk, M. D., & LeDoux, J. E. (2002). Parallels between cerebellum-and amygdala-dependent conditioning. *Nature Reviews Neuroscience*, 3(2), 122–131.
<https://doi.org/10.1038/nrn728> (<https://doi.org/10.1038/nrn728>)
- Musazzi, L., Tornese, P., Sala, N., & Popoli, M. (2017). Acute or chronic? A stressful question. *Trends in Neurosciences*.
<https://doi.org/10.1016/j.tins.2017.07.002>
(<https://doi.org/10.1016/j.tins.2017.07.002>)
- Pellman, B. A., & Kim, J. J. (2016). What can ethobehavioral studies tell us about the brain's fear system? *Trends in Neurosciences*, 39(6), 420–431. <https://doi.org/10.1016/j.tins.2016.04.001>
(<https://doi.org/10.1016/j.tins.2016.04.001>)
- Schmaal, L., Veltman, D. J., Erp, T. G. M. van, Sämann, P. G., Frodl, T., Jahanshad, N., ... Hibar, D. P. (2016). Subcortical brain alterations in major depressive disorder: Findings from the ENIGMA major depressive disorder working group. *Molecular Psychiatry*, 21(6), 806–812. <https://doi.org/10.1038/mp.2015.69>
(<https://doi.org/10.1038/mp.2015.69>)