

IN BRIEF...

BAC-driven miRNA gene expression knockdown

KA Garbett, S Horvath, PJ Ebert, MJ Schmidt, K Lwin, **A Mitchell**, P Levitt and K Mirnics (2010). Novel animal models for studying complex brain disorders: BAC-driven miRNA-mediated *in vivo* silencing of gene expression. *Mol. Psychiatry*. doi: 10.1038/mp.2010.1

Animal models of disease represent one of the most powerful methods of analyzing the pathophysiological mechanisms of genetic disorders. However, the development of such models is often time-consuming, complex, and carries nonspecific caveats, such as the imprecise deletion of a gene of choice. Using bacterial artificial chromosomes, cell-type specific promoters, a standard reporter, and a microRNA mechanism for gene silencing, Garbett *et al.* present a powerful mechanism to specifically reduce gene expression *in vivo*. As microRNAs are of small size, they anticipate that this new method could simultaneously silence multiple genes in a cell-type specific manner. Accordingly, these transgenic mice would allow exquisite precision in determining the effects of a given set of genes on the presentation of disease.

Pain Pathways: Neuropeptide Y may be targeted to relieve pain

RG Wiley, **LL Lemons** and RH Kline IV (2009). Neuropeptide Y receptor-expressing dorsal horn neurons: role in nocifensive reflex responses to heat and formalin. *Neuroscience*. **161** (1): 139-147.

There is an endless list of reasons why individuals seek treatment for pain, but the molecular mechanisms that underlie pain perception are unclear. Wiley *et al.* demonstrate how Neuropeptide Y (NPY) and its receptor Y1 (Y1R) function in the rodent spinal cord to mediate nociception. After intrathecal injection of saporin toxin conjugated to NPY to selectively kill Y1R-expressing neurons in the dorsal horn of the spinal cord, rats displayed an increased latency to withdraw their paws from noxious hot stimuli. The rodents also had a significant decrease in nocifensive behavior when presented with the hot stimuli or when injected with formalin in the plantar region of the hind paw, as measured by licking and guarding events. This toxin-based approach allows researchers to selectively examine groups of neurons involved in the perception of pain and tease apart each group's contribution. These studies could prove to have a significant impact on the field of pain research and may provide researchers with some insight into alternative approaches to treat pain.

What's unusual about that? Neural substrates for the detection of novel, unusual stimuli

Blackford JU, Buckholz JW, **Avery SN**, Zald DH (2010). A unique role for the human amygdala in novelty detection. *Neuroimage*. **50**(3):1188-93.

Novelty detection is an important trait in perceiving and responding to our environment. In particular novel, yet unusual or uncommon stimuli that are behaviorally salient can engage specific neural mechanisms involved in emotional learning and memory. In this study, the authors used functional magnetic resonance imaging to observe blood oxygen level dependent (BOLD) responses in the human amygdala and hippocampus when they presented participants with novel, common stimuli (e.g., chair, clock, tree) versus novel, unusual stimuli (e.g., Prague Dancing House, futuristic skyscraper, leafy sea dragon). Blackford and colleagues found that novel, common stimuli showed robust BOLD activation in both the amygdala and the hippocampus. However, only the amygdala showed a preferential activation for the novel, unusual stimuli, compared to the novel common stimuli. These results lead the authors to speculate that within the novelty detection circuit, the amygdala plays a distinct role in uniquely responding to a specific category of stimuli, namely those that are novel and unusual.

change or neural reorganization. Powers and colleagues found significant effects in the temporal dynamics of the multisensory binding window after only one day of training, with stable effects observed even after one week. This alludes to the fact that the brain is capable of adapting quickly to new environmental situations and long term memory consolidation may play a role in strengthening learned traits. Thus results from this study would support the idea that the pairing of a sensory stimulus with behavioral salience (like feedback in this study) is crucial for sensory reorganization of adult cortical space.

It is intriguing to consider where and how the plasticity associated with this temporal binding window is modulated. In humans, recent neuroimaging studies reported a large, dynamic network of areas including the insular cortex, posterior parietal and superior temporal cortices, all critically involved in the perception of audiovisual stimuli. The neuronal oscillations among different cortical populations have also been shown to play a potential role in multisensory processing and temporal binding. At a more cellular level, the temporal tuning profile of multisensory neurons has been associated with adult plasticity in various sensory systems, with basal cholinergic signals acting as an instructive cue. This indicates that the synchronous role of both cortical and subcortical mechanisms may be responsible for temporal plasticity in multisensory systems.

The present work by Powers and colleagues and future extensions of similar studies holds particular clinical significance, especially in designing tailored intervention strategies for disorders such as dyslexia, autism and schizophrenia where altered multisensory temporal processes have been observed.

Original Research Article:

AR Powers III, AR Hillock and MT Wallace (2009). Perceptual training narrows the temporal window of multisensory binding. *J Neurosci*. **29** (39): 12265-12274.

Development: birth, life, death, cleanup, repeat.

The development of a nervous system entails several obvious processes such as the proliferation of cells, the elaboration of dendrites, or the wiring of functional axonal circuits, yet it is now becoming clear that the less publicized (and slightly more sinister) mechanisms of programmed cell death and debris clearance are a vital component of nervous growth. For example, in the mouse dorsal root ganglia (DRG) over 50% of sensory neurons undergo