Washington University School of Medicine Digital Commons@Becker

Presentations

2006: Alcohol and Tobacco Dependence: from Bench to Bedside

2006

Addiction: Reward, motivation and stress

George F. Koob Scripps Research Institute

Follow this and additional works at: http://digitalcommons.wustl.edu/guzepresentation2006 Part of the <u>Medicine and Health Sciences Commons</u>

Recommended Citation

Koob, George F., "Addiction: Reward, motivation and stress" (2006). *Presentations*. Paper 2 Samuel B. Guze Symposium on Alcoholism. http://digitalcommons.wustl.edu/guzepresentation2006/2

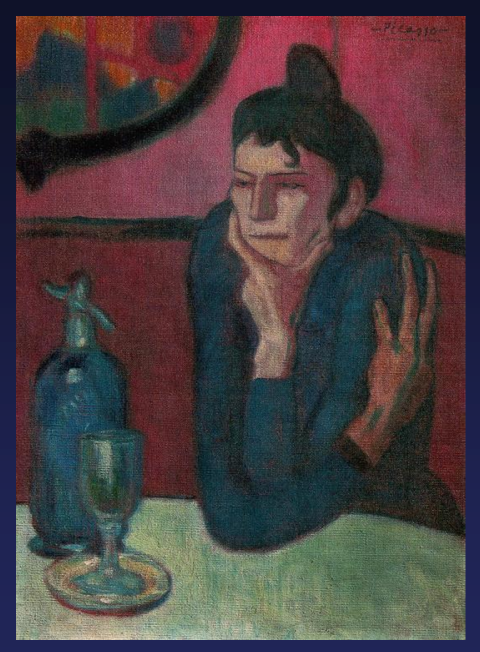
This Presentation is brought to you for free and open access by the 2006: Alcohol and Tobacco Dependence: from Bench to Bedside at Digital Commons@Becker. It has been accepted for inclusion in Presentations by an authorized administrator of Digital Commons@Becker. For more information, please contact engeszer@wustl.edu.

Addiction: Reward, Motivation and Stress

George F. Koob, Ph.D. Professor Molecular and Integrative Neurosciences Department The Scripps Research Institute La Jolla, California

The Neurocircuitry of Drug Addiction: Neuroadaptive Mechanisms from the "Dark Side"

- What is Addiction?
 - **1.** Conceptual framework
 - 2. The 'dark side' of compulsivity
- Animal Models for the Motivational Effects of Dependence
 - **1. Brain stimulation reward**
 - 2. Place aversion
 - 3. Anxiogenic-like responses in the plus maze and defensive burying tests
 - 4. Escalation in drug self-administration with prolonged access
- A Role for Corticotropin-Releasing Factor in Drug Addiction
 - 1. Cocaine
 - 2. Nicotine
 - 3. Heroin
 - 4. Alcohol
- Future Directions
 - **1.** Development of CRF₁ antagonists for treatment of addiction
 - 2. The neurocircutry of emotional behavior



"Absinthe Drinker" Pablo Picasso (1910)

Key Definitions

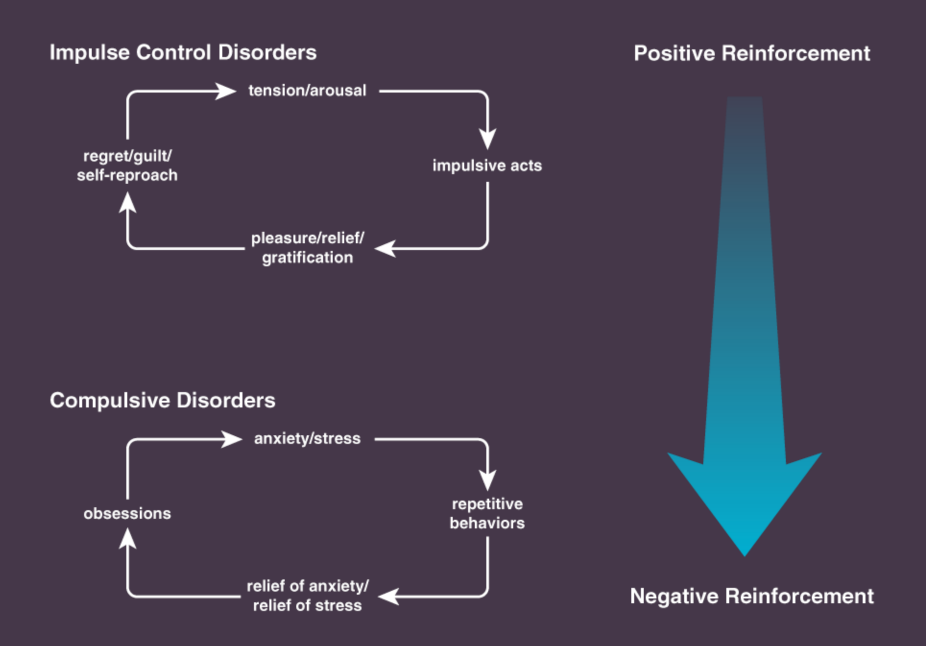
Drug Addiction — Chronically relapsing disorder that is characterized by a compulsion to seek and take drug, loss of control in limiting intake, and emergence of a negative emotional state (e.g. dysphoria, anxiety, irritability) when access to the drug is prevented (here, defined as the "dark side" of addiction)

Extended Amygdala — Forebrain macrostructure composed of central medial amygdala, bed nucleus of the stria terminalis, and a transition zone in the medial part of the nucleus accumbens

Corticotropin-Releasing Factor — 41 amino acid polypeptide "brain stress" neurotransmitter that controls hormonal, sympathetic, and behavioral responses to stressors

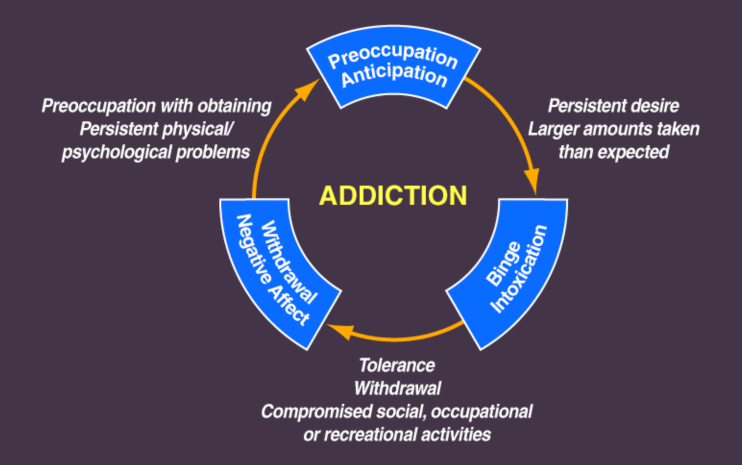
Drug Addiction

Drug addiction is conceptualized as a chronic relapsing syndrome that moves from an impulse control disorder involving positive reinforcement to a compulsive disorder involving negative reinforcement



From: Koob GF, Alcohol Clin Exp Res, 2003, 27:232-243.

Stages of the Addiction Cycle



Animal Models for the Motivational Components of Dependence

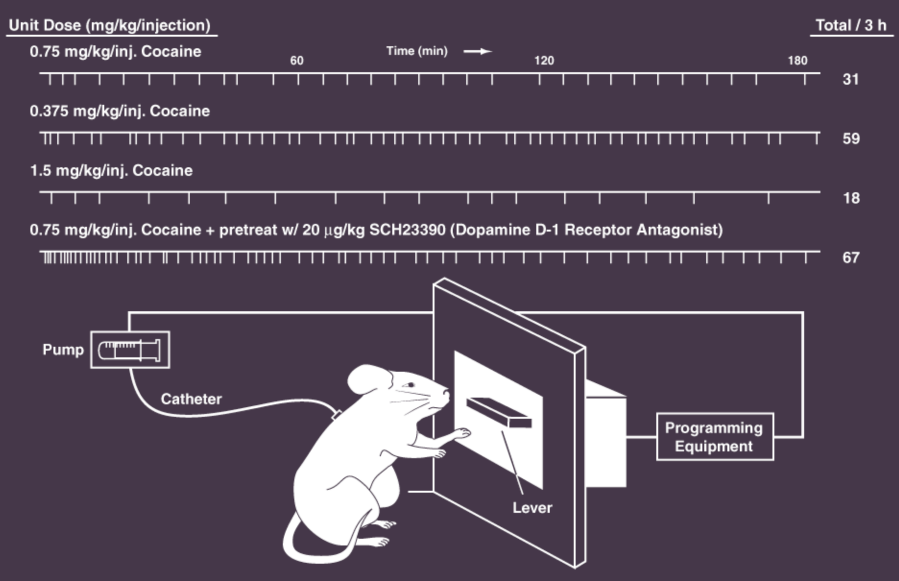
Animal Models for the Withdrawal/Negative Affect Stage

- **1. Brain stimulation reward**
- 2. Place aversion
- **3.** Anxiogenic-like responses in elevated plus maze and defensive burying

Animal Models for the Transition to Addiction

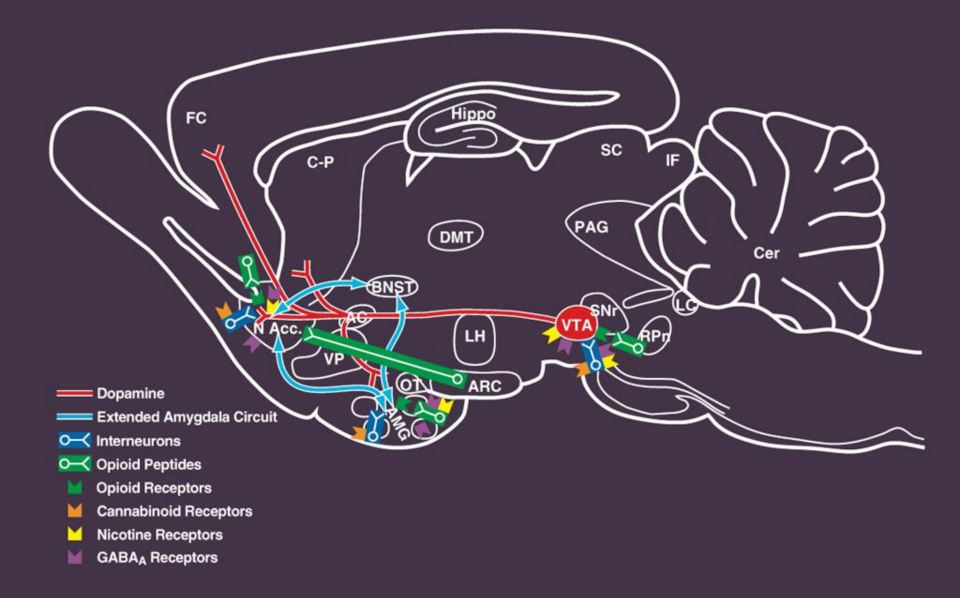
- **1. Drug taking in selected lines of drug preferring animals**
- 2. Withdrawal-induced drug taking
- **3.** Escalation in drug self-administration with prolonged access
- 4. Drug taking despite aversive consequences

Cocaine Self-Administration

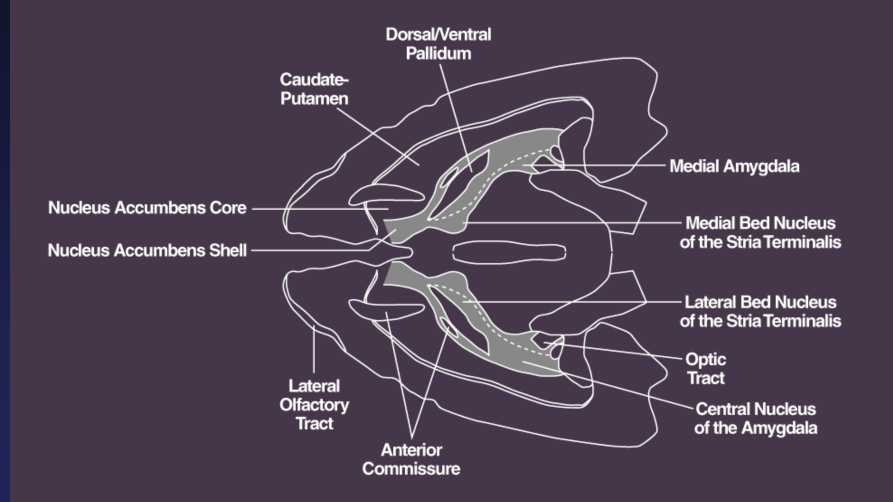


From: Caine SB, Lintz R and Koob GF. in Sahgal A (ed) <u>Behavioural Neuroscience: A Practical Approach</u>, vol. 2, IRL Press, Oxford, 1993, pp. 117-143.

Neurochemical Circuitry in Drug Reward

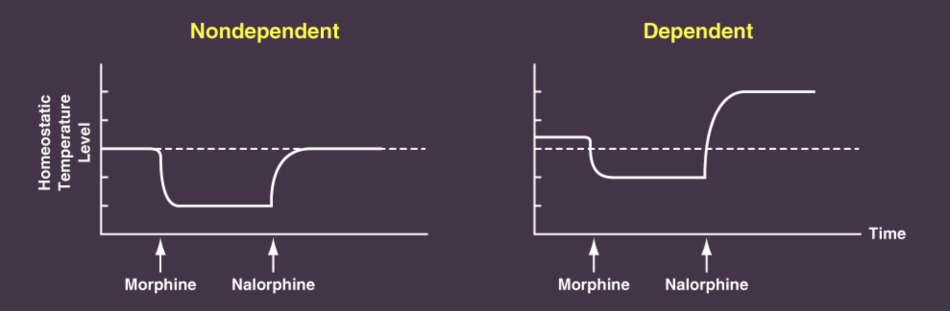


Potential Substrates in the Extended Amygdala for the Motivational Effects of Drug Dependence



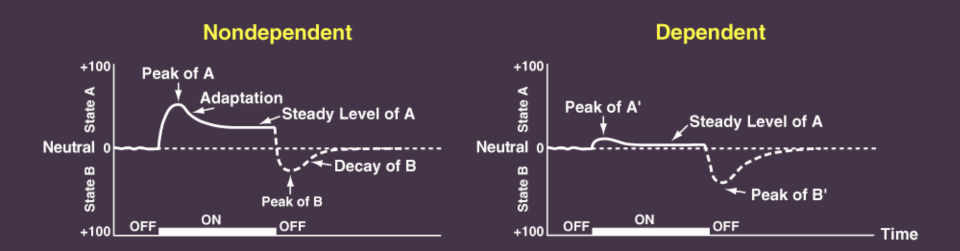
Modified from: Heimer L and Alheid G, Piecing together the puzzle of basal forebrain anatomy. In: Napier TC, Kalivas PW and Hanin I (Eds), <u>The Basal Forebrain: Anatomy to Function</u> (series title: <u>Advances in Experimental</u> <u>Medicine and Biology</u>, Vol. 295), Plenum Press, New York, 1991, pp. 1-42.

Equilibrium State for a Homeostatic Regulatory System in a Nondependent and Dependent Organism



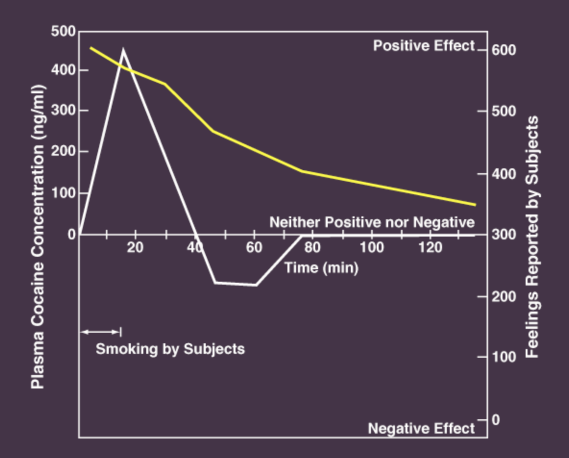
From: Martin WR, A homeostatic and redundancy theory of tolerance to and dependence on narcotic analgesics. in Wikler A (Ed.), <u>The Addictive States</u> (series title: *Its Research Publications*, vol. 46), Williams and Wilkins, Baltimore, 1968, pp. 206-225.

Standard Pattern of Affective Dynamics Produced by Novel and Repeated Unconditioned Stimulus



From: Solomon RL, American Psychologist, 1980, 35:691-712.

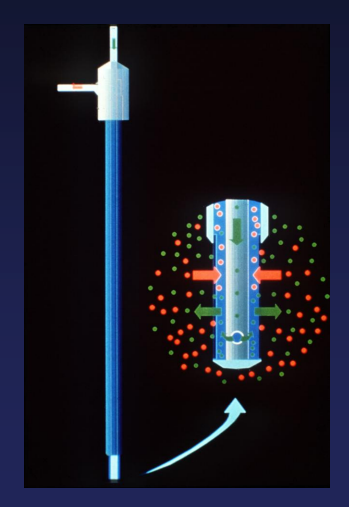
Mood Changes Associated with Plasma Levels of Cocaine During Coca Paste Smoking



Dysphoric Feelings followed the initial euphoria in experimental subjects who smoked cocaine paste, even though the concentration of cocaine in the plasma of the blood remained relatively high. The dysphoria is characterized by anxiety, depression, fatigue and a desire for more cocaine.

From: Van Dyke C and Byck R, Cocaine, Scientific American, 1982, 246:123-141.

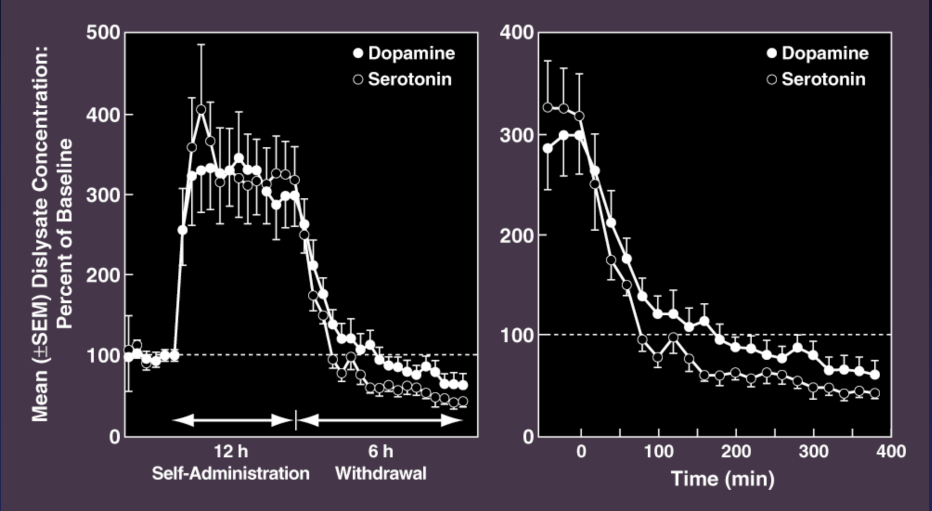
Sampling of Interstitial Neurochemicals by *in vivo* Microdialysis



- Allows sampling of neurochemicals in conscious animals (correlate brain chemistry with behavior).
- Implanted so that semi-permeable probe tip is in specific brain region of interest.
- Substances below the membrane MW cutoff diffuse across membrane based on concentration gradient.
- Both neurochemical sampling and localized drug delivery are possible.

Collaborators: Dr. Friedbert Weiss, Dr. Larry Parsons, Dr. Emilio Merlo-Pich, Dr. Regina Richter

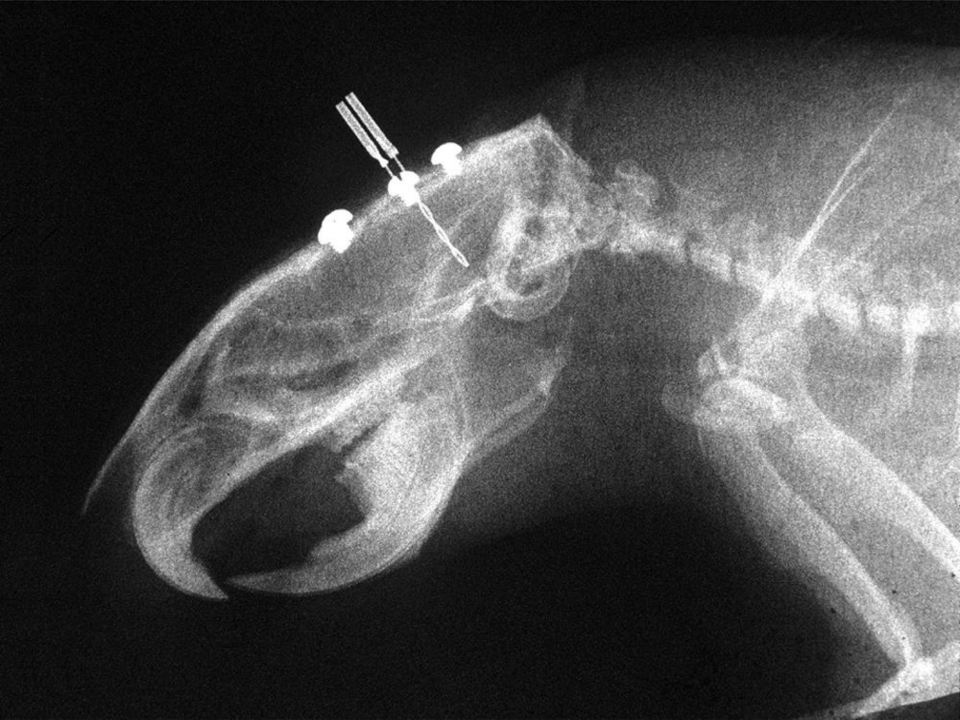
Extracellular DA and 5-HT in the Nucleus Accumbens During Cocaine Self-Administration and Withdrawal



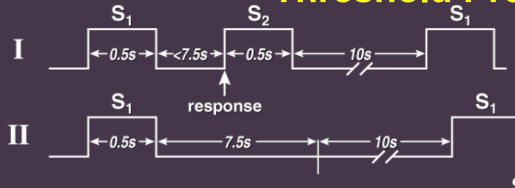
From: Parsons LH, Koob GF and Weiss F, <u>J Pharmacol Exp Ther</u>, 1995, 274:1182-1191.

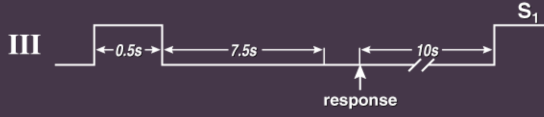
Drug Withdrawal

Withdrawal from chronic drugs of abuse produces a reward (motivational) dysregulation as measured by thresholds for intracranial self-stimulation



Intracranial Self-Stimulation (ICSS) Threshold Procedure





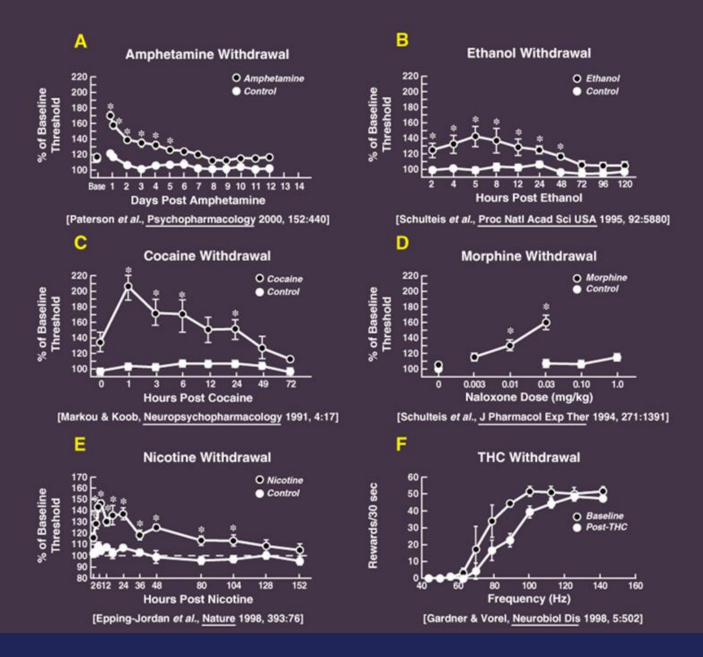
Current

(μΑ)	Descending			Ascending			Descending			Ascending			
180	+	+	+				+	+	+	+	+	+	
175	+	+	+	+	+	+	+	+	+	+	+	+	
170	+	+	+	+	+	+	+	+	+	+	+	+	
165	+	-	+	-	-	-	+	-	-	-	-	-	
160	-	-	-	-	+	-	-	+	+	-	-	-	
155	-	-	-	-	-	-	-	-	-	-	-	+	
150				-	-	-	-	-	-	-	-	-	
145										-	-	-	
	162.5		167.5			157.5			167.5				
	Threshold = 163.75 μA												



Adapted from: Markou A and Koob GF, Physiol Behav, 1992, 51:111-119.

Elevations in ICSS Reward Thresholds During Withdrawal



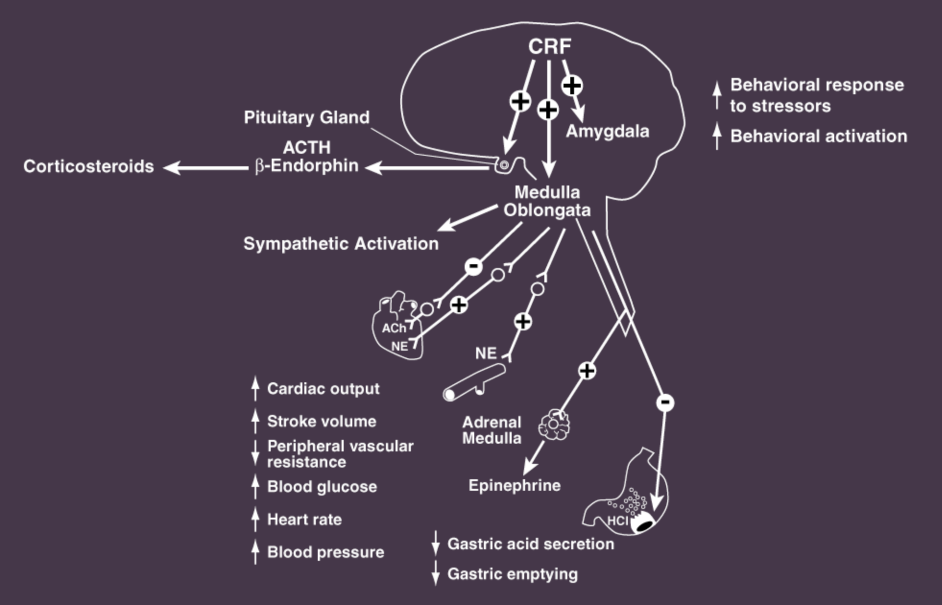
Reward Transmitters Implicated in the Motivational Effects of Drugs of Abuse

Positive Hedonic Effects

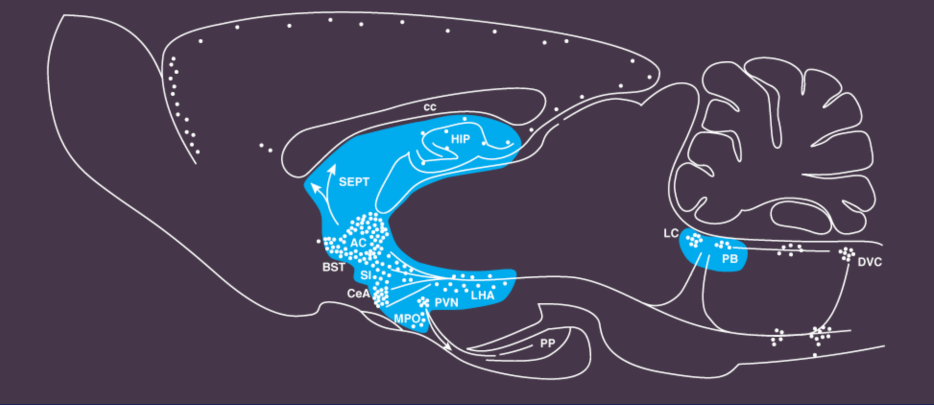
- **†** Dopamine
- **†** Opioid peptides
- Serotonin
- **†** GABA

Negative Hedonic Effects of Withdrawal
Dopamine ... "dysphoria"
Opioid peptides ... pain
Serotonin ... "dysphoria"
GABA ... anxiety, panic attacks

CNS Actions of Corticotropin-Releasing Factor (CRF)



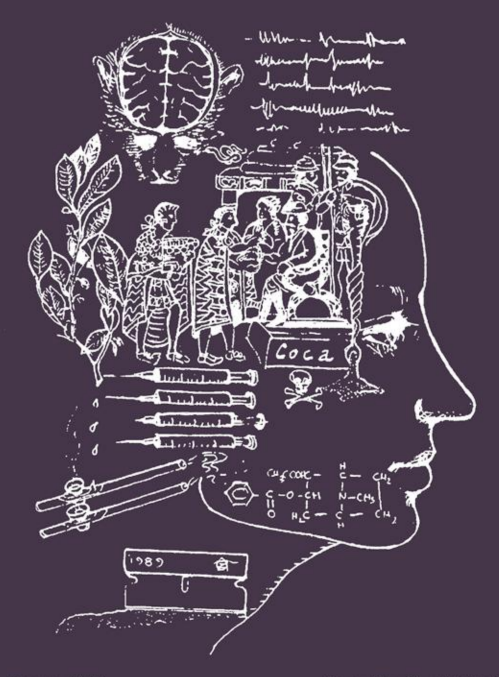
Major CRF-Immunoreactive Cell Groups and Fiber Systems in the Rat Brain



From: Swanson LW, Sawchenko PE, Rivier J and Vale W, <u>Neuroendocrinology</u>, 1983, 36:165-186.

CRF Produces Arousal, Stress-like Responses, and a Dysphoric, Aversive State

Paradigm	CRF Agonist	CRF Antagonist
Acoustic startle	Facilitates startle	Blocks fear-potentiated startle
Elevated plus maze	Suppresses exploration	Reverses suppression of exploration
Defensive burying	Enhances burying	Reduces burying
Fear conditioning	Induces conditioned fear	Blocks acquisition of conditioned fear
Cued electric shock	Enhances freezing	Attenuates freezing
Taste / Place Conditioning	Produces place aversion	Weakens drug-induced place aversion





Chronic cocaine administration produces a dependence syndrome that is reversed by blockade of CRF function.

Defensive Burying: Active Anxiety-Like Behavior





Habituation

- Two 45-min sessions in test cage
- No shock probe present

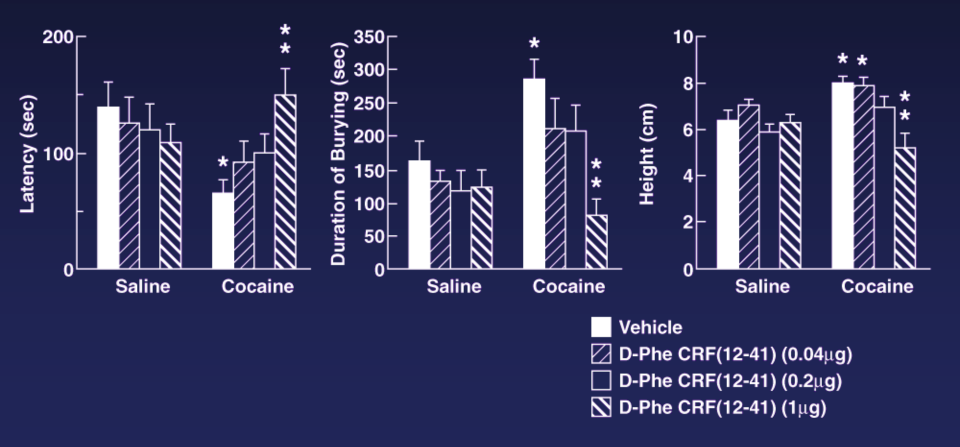
Testing

- Electrified shock probe present
- Probe delivers a single, < 1 sec, 1.5 mA shock on contact
- Probe is shut off after shock
- Defensive burying scored for 10 min

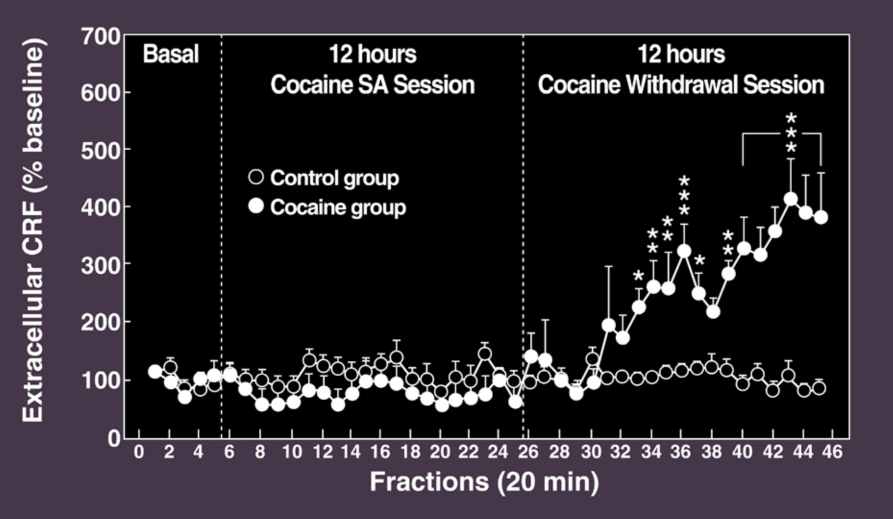
Endpoints

- Latency to bury
- Duration of burying
- Duration of other active behaviors

Effect of CRF Antagonist D-Phe-CRF₁₂₋₄₁ Administered ICV on Anxiogenic-Like Effect Following Chronic Cocaine Administration



From: Basso AM, Spina M, Rivier J, Vale W and Koob GF, <u>Psychopharmacology</u>, 1999, 145:21-30.



Protocol for Drug Escalation

1) Initial Training Phase

All Rats (n=24): 2-hr SA session Fixed Ratio 1 0.25 mg cocaine/injection 2) Escalation Phase

Short Access (n=12) 22 x 1-hr SA session

Long Access (n=12) 22 x 6-hr SA session 3) Testing Phase

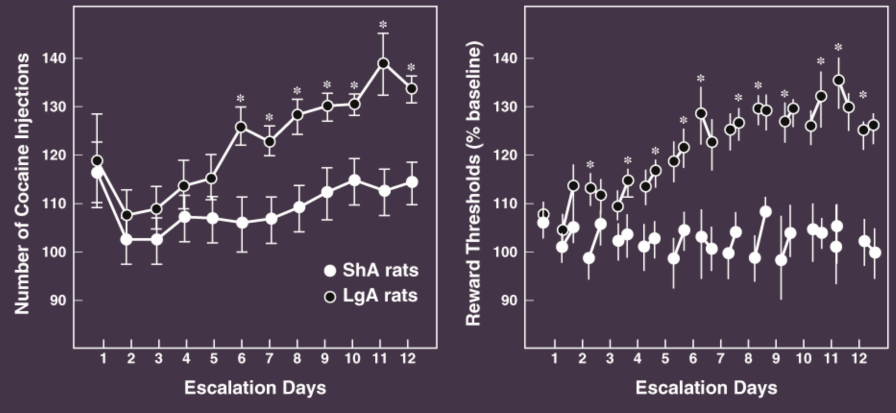
Dose-response for neuropharmacological probes

Protocol from: Ahmed SH and Koob, Science, 1998, 282:298-300.

Change in Brain Stimulation Reward Thresholds in Long-Access (Escalation) vs. Short-Access (Non-Escalation) Rats

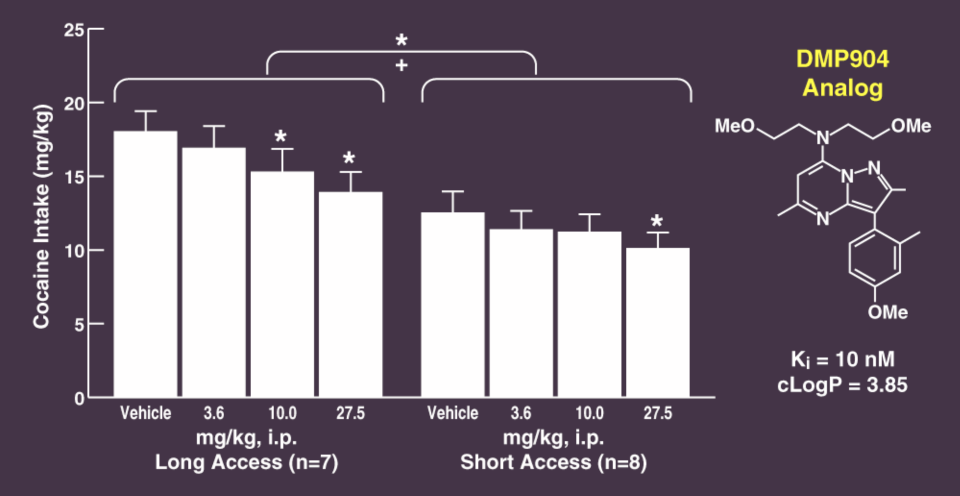


Brain Stimulation Reward Thresholds

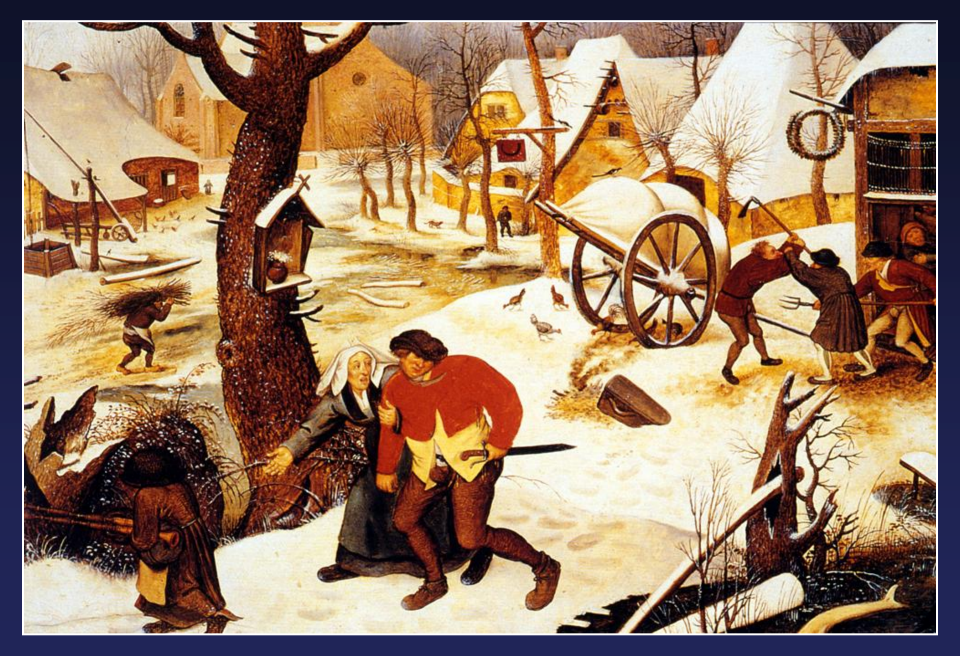


From: Ahmed SH, Kenny PJ, Koob GF and Markou A, Nature Neurosci, 2002, 5:625-627.

Dose-Dependent Decrease of Cocaine Intake with Administration of a CRF₁ Antagonist



From: Specio SE, Zorrilla EP, O'Dell LE and Koob GF, unpublished results.



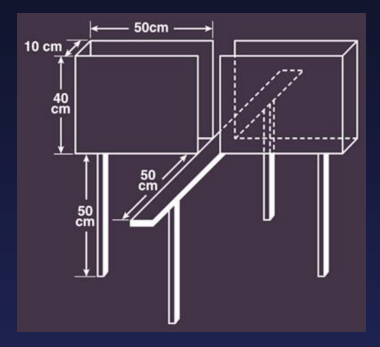
Pieter Bruegel

Alcohol

Chronic alcohol exposure produces a dependence syndrome that is reversed by blockade of CRF function.

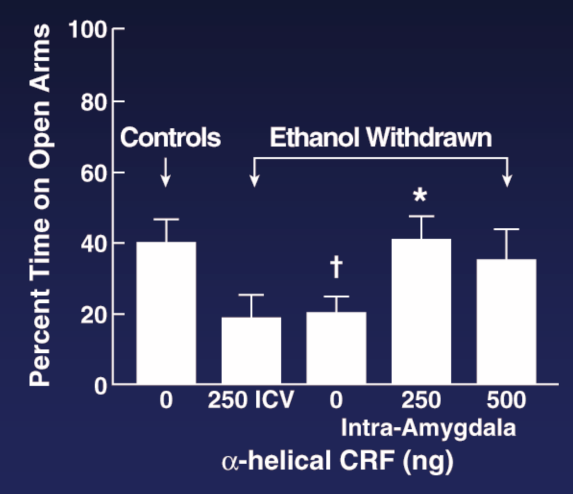
Elevated Plus Maze





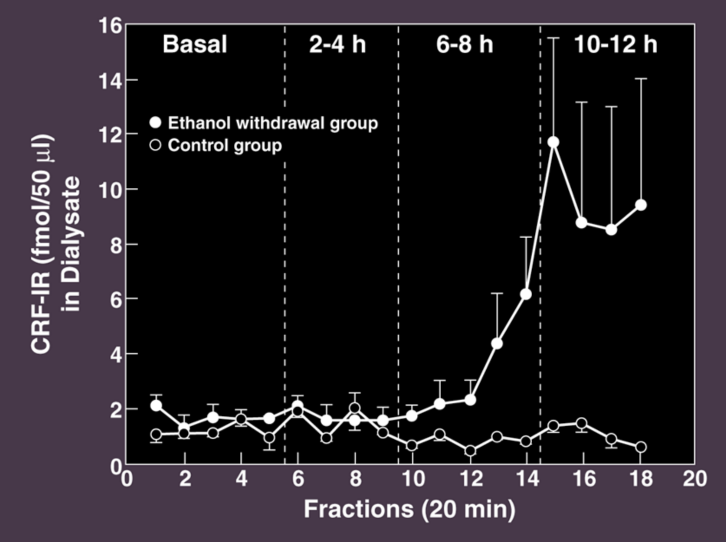
- Unconditioned approach/avoidance behavior
- 3 underlying factors: anxiety, activity, assessment of risk
- Predictive validity for anxiolytic and anxiogenic drugs

Competitive CRF Antagonist α-Helical CRF₉₋₄₁ Injected into Central Nucleus of the Amygdala Blocks the Anxiogenic Effects of Alcohol Withdrawal



From: Rassnick S, Heinrichs SC, Britton K and Koob GF, Brain Res, 1993, 605:25-32.

Extracellular CRF Levels in the Central Amygdala During Ethanol Withdrawal



From: Merlo-Pich E, Lorang M, Yeganeh M, Rodriquez de Fonseca F, Koob GF and Weiss F, <u>J Neurosci</u>, 1995, 15:5439-5447.



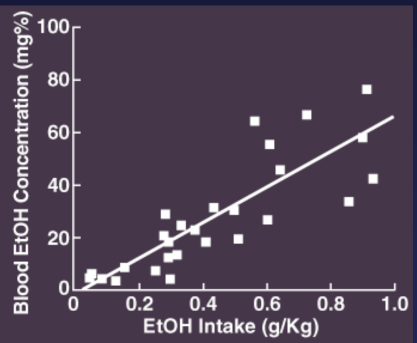
Protocol for Initiation of Lever Pressing for Oral Ethanol Self-Administration in the Rat

Training		Saccharin (w/v)	EtOH (w/v)
Days	1-3	0.2%	0% *
Days	4-9	0.2%	5% *
Day	10	-	5% *
Days	11-12	0.2%	5%
Day	13	-	5%
Day	14	0.2%	8%
Days	15-16	-	8%
Day	7	0.2%	10%
Day	18+	-	10% *

Rats trained to lever press on a FR-1 schedule

Ethanol added to the saccharin solution

Access to ethanol and water or ethanol + saccharin and water

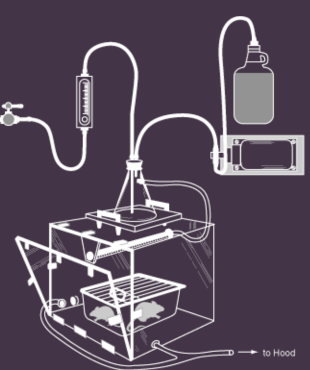


Initiation of the free-choice operant task: ethanol (10%) and water

From: Rassnick S, Pulvirenti L and Koob GF, Alcohol, 1993, 10:127-132.

Ethanol Dependence Induction

Ethanol Vapor Chambers



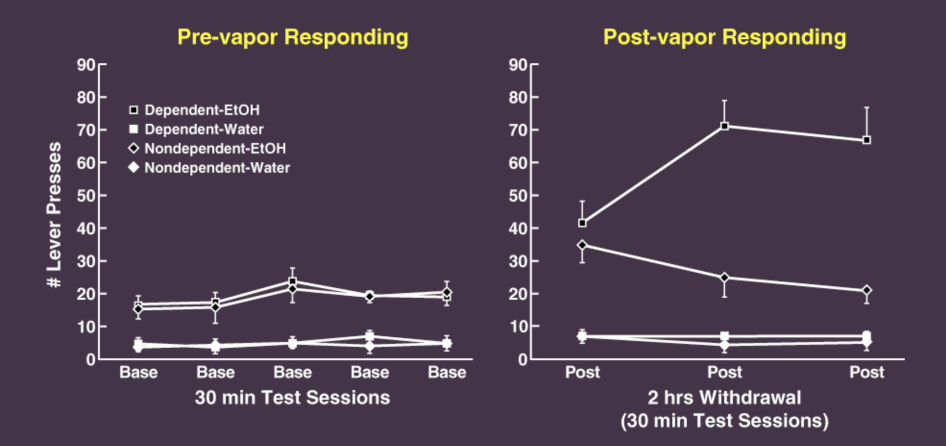
- Ethanol vapor concentrations range from 22-27 mg/liter
- BAL's are determined every 3 days and ethanol flow is adjusted to maintain BAL's of 150-200 mg%
- Dependence is reliably induced following 2 weeks of exposure
- Control rats are placed in identical chambers into which only air is pumped

Ethanol Liquid Diet

- 8.7% (w/v) ethanol with 35% ethanol-derived calories
- Consists of ethyl alcohol, chocolate flavored sustacal, vitamin and mineral diet fortification
- With unlimited access, maintains BALs over 140 mg%
- Dependence is reliably induced following 2 week exposure
- Control rats are fed liquid diet substituting sucrose for ethanol

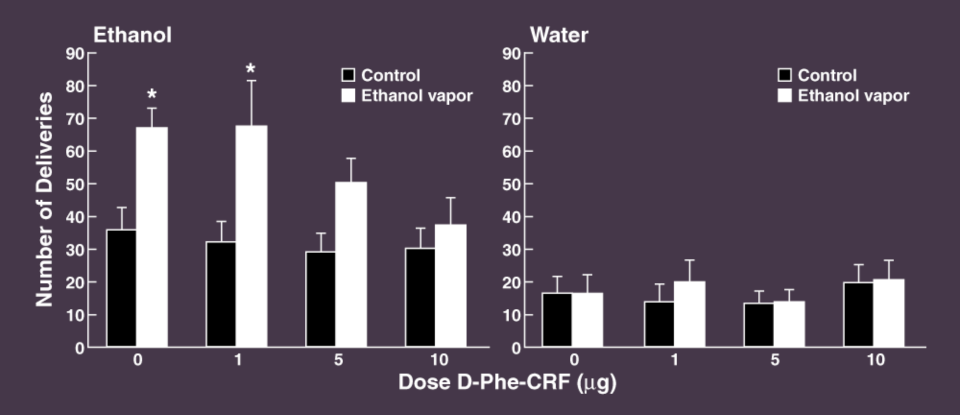


Enhanced Ethanol Self-Administration During Withdrawal in Dependent Animals



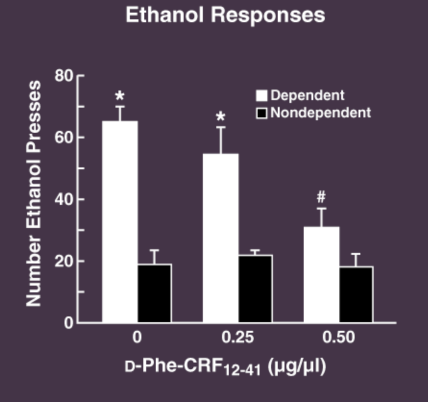
Effects of a Competitive CRF Antagonist Injected ICV on Ethanol Self-Administration During Withdrawal in Dependent Rats

(60 min session 2 h into withdrawal)



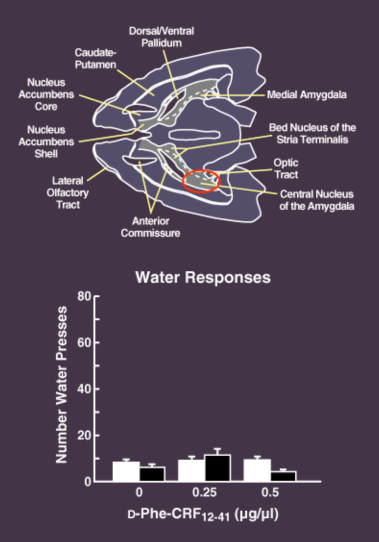
From: Valdez GR, Roberts AJ, Chan K, Davis H, Brennan M, Zorrilla EP and Koob GF, <u>Alcohol Clin Exp Res</u>, 2002, 26:1494-1501.

Effect of CRF Antagonist D-Phe-CRF₁₂₋₄₁ – Central Nucleus of the Amygdala –

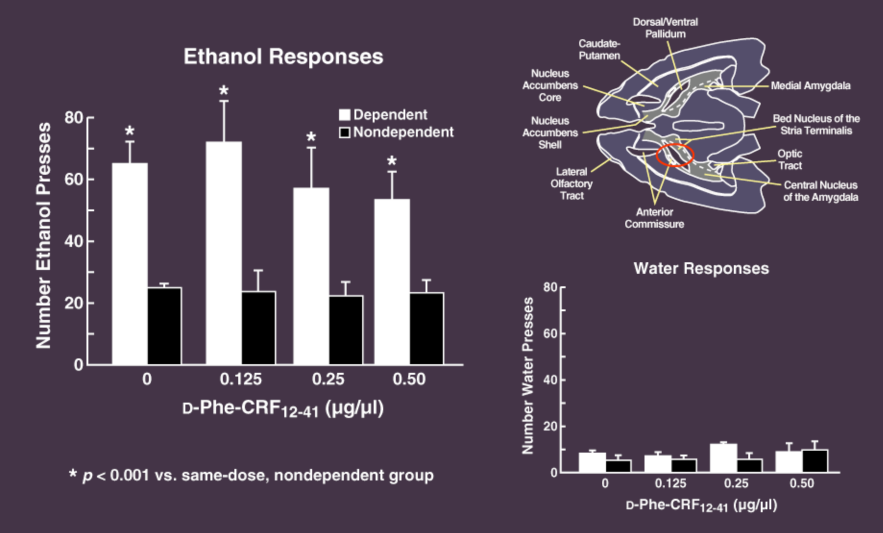


* p < 0.001 vs. same-dose, nondependent group # p < 0.001 vs. dependent, vehicle group

From: Funk C, O'Dell LE and Koob GF, unpublished results.

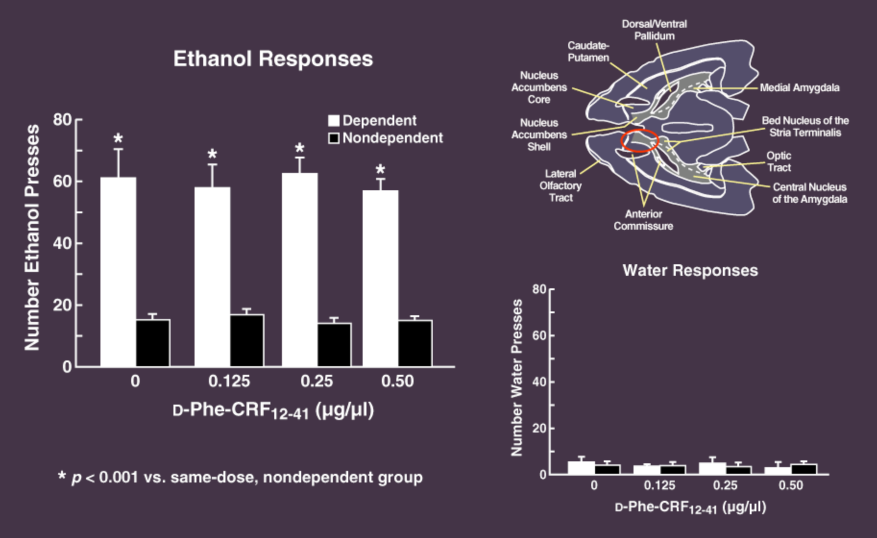


Effect of CRF Antagonist D-Phe-CRF₁₂₋₄₁ – Lateral Bed Nucleus of the Stria Terminalis –



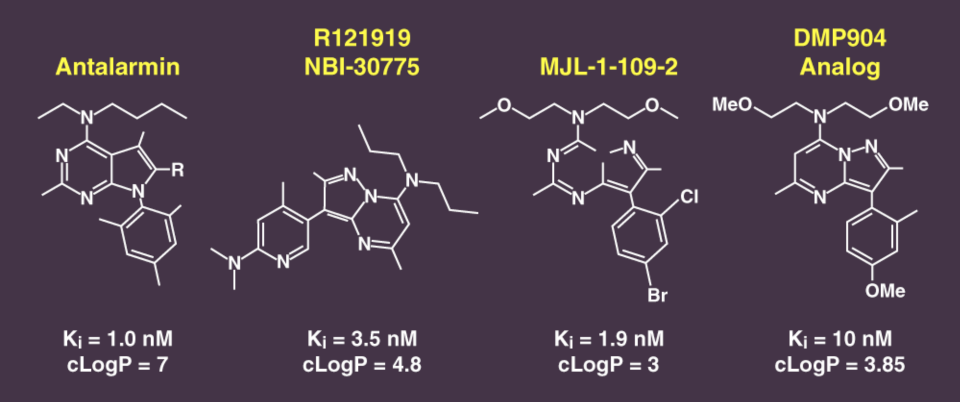
From: Funk C, O'Dell LE and Koob GF, unpublished results.

Effect of CRF Antagonist D-Phe-CRF₁₂₋₄₁ – Nucleus Accumbens Shell –

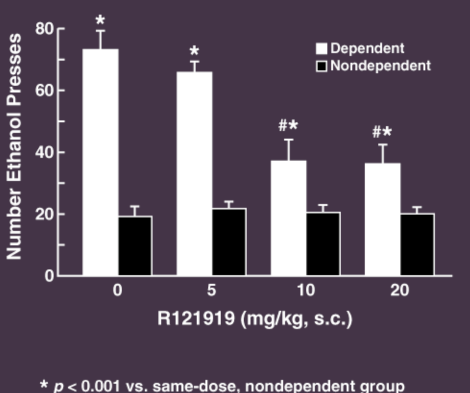


From: Funk C, O'Dell LE and Koob GF, unpublished results.

CRF₁ Specific Antagonists



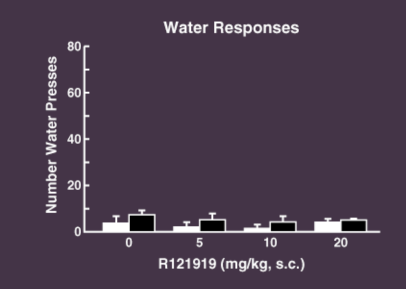
CRF₁ Specific Antagonists R121919



Ethanol Responses

* p < 0.001 vs. same-dose, nondependent group # p < 0.001 vs. dependent, vehicle group</p>

- administered s.c.
 60 min pre-incubation
- n = 9
- HBC (20% w/v)



N $K_i = 3.5 \text{ nM}$



From: Funk C, Zorrilla EP, Lee MJ, Rice KC and Koob GF, unpublished results.

Interaction of CRF Antagonists in Animal Models of Protracted Abstinence

1. CRF antagonists injected into the extended amygdala block stress-induced reinstatement of drug seeking

Erb S, Salmaso N, Rodaros D and Stewart J, Psychopharmacology, 2001, 158:360-365 Liu X and Weiss F, J Neurosci, 2002, 22:7856-7861 Funk D, Li Z, Shaham Y and Le AD, Neuroscience, 2003, 122:1-4

2. CRF antagonists injected i.c.v. block stress-induced anxiogenic-like responses and excessive drinking during protracted abstinence

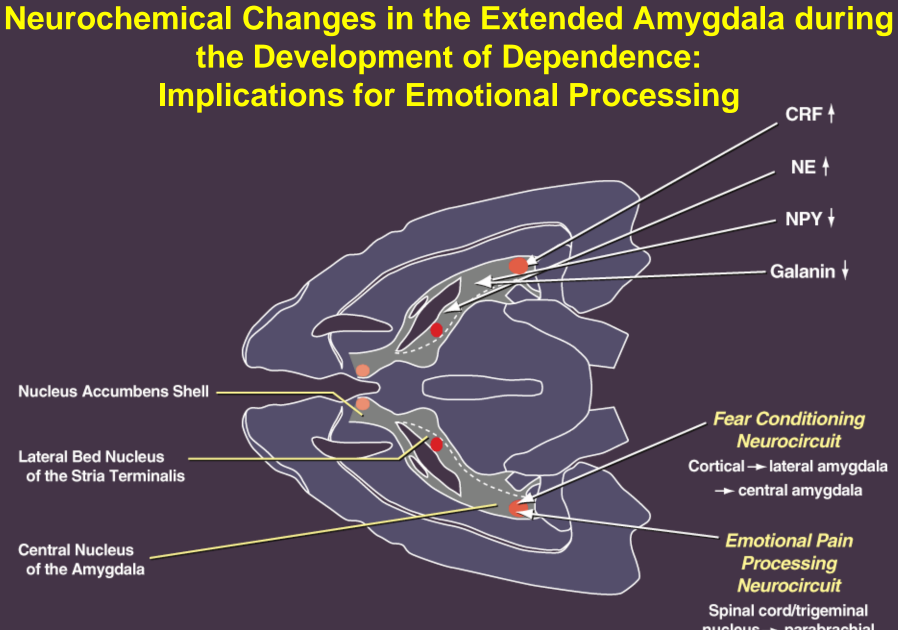
Valdez GR, Zorrilla EP, Roberts AJ and Koob GF, Alcohol, 2003, 29:55-60.

3. CRF₁ knockout mice show a blunted anxiogenic-like response to alcohol withdrawal and a blockade of excessive drinking during protracted abstinence

Chu K, Koob GF, Cole M and Roberts AJ, submitted

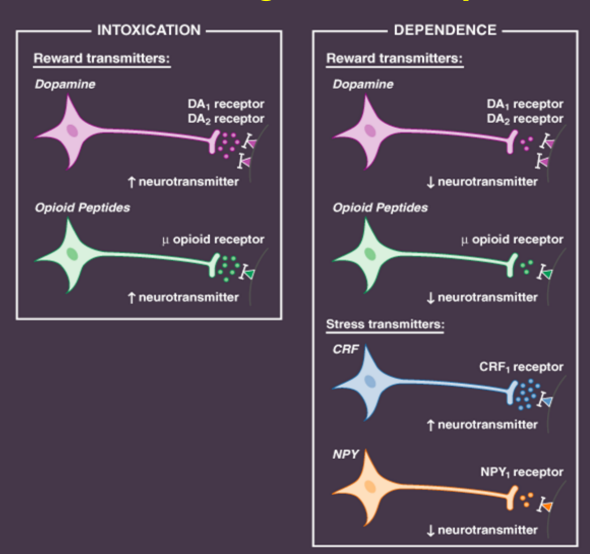
Extended Amygdala

The extended amygdala is a rich substrate for neurochemical and neurocircuitry interactions that produce the "dark side" of motivation.



nucleus → parabrachial complex → central amygdala

Neurochemical Changes Associated with the Transition from Drug Use to Dependence



From: Roberts AJ and Koob GF, Alcohol: ethanol antagonists/amethystic agents. in Adelman G and Smith BH (Eds.), Encyclopedia of Neuroscience, 3rd edn, Elsevier, New York, 2003 [http://203.200.24.140:8080/Neuroscience].

Conclusions

CRF in the extended amygdala is recruited during the development of dependence and has motivational significance for drug seeking.

Compulsive drug taking associated with addiction derives both from decreases in reward neurotransmission and from recruitment of antireward systems ("dark side" of addiction).

Other neurochemical elements in the extended amygdala—such as norepinephrine, NPY and galanin—may have a role in motivational neuroadaptation associated with drug dependence.

The common interface in the extended amygdala of the neurochemistry of addiction and pain and fear conditioning pathways provides a heuristic framework for exploring the neural basis of negative emotional states.

Neurobiology of Drug Addiction Koob Laboratory

Post-Doctoral Fellows Cindy Funk Brendan Walker Tom Greenwell Sandy Ghozland Nick Gilpin Chitra Mandyam Sunmee Wee Research Assistants Bob Lintz Richard Schroeder Elena Crawford Molly Brennan Maury Cole Tess Kimber Yanabel Grant Ron Smith

Special thanks to: Mike Arends (Senior Research Assistant)

Janet Hightower (Biomedical Graphics Dept) Administrative Assistants Lisa Maturin Mellany Santos Marisa Gallego

Neurobiology of Drug Addiction Current Collaborators

-loyd Bloom	Jean Rivier
Barbara Mason	Catherine Rivier
Vichel Le Moal	Tamas Bartfai
₋uis Stinus	George Siggins
Friedbert Weiss	Marisa Roberto
Athina Markou	Kenner Rice
Amanda Roberts	Kim Janda
_arry Parsons	Laura O'Dell
Pietro Sanna	Robert Purdy
₋uigi Pulvirenti	Walter Francescon
Eric Zorrilla	Sheila Specio
Nylie Vale	Marc Azar

Support from:

National Institute on Alcohol Abuse and Alcoholism National Institute on Drug Abuse National Institute of Diabetes and Digestive and Kidney Diseases Pearson Center for Alcoholism and Addiction Research