Psychoactive Drugs and Addiction

What are psychoactive drugs? Substances that influence subjective experience and behavior by acting on the nervous system.

What are some of the most common psychoactive drugs?
1. Sedative hypnotics and anxiolytics (antianxiety drugs)
2. Antipsychotic drugs
3. Antidepressant drugs
4. Opiates (analgesics)
5. Stimulants

I. Sedative hypnotics and anxiolytics:
   Most common:
   1. Alcohol
   2. Barbiturates (ex. pentobarbital, sodium amytal – truth serum)
   3. Benzodiazepines (ex., valium™, librium™)

   Also known as “mild tranquilizers”

How do sedative hypnotics and antianxiety drugs work? They bind to the GABA \(_A\) receptor complex; alcohol and barbiturates act just like endogenous GABA and open the chloride channel directly; benzodiazepines facilitate the action of endogenous GABA, but does not directly open chloride channels.

II. Antipsychotic drugs: used to treat psychotic conditions such as schizophrenia, paranoia, etc.
   Most common:
   1. Phenothiazines (ex., chlorpromazine - Thorazine™)
   2. Butyrophenones (ex., haloperidol - Haldol™)

   Also known as “major tranquilizers”

How do antipsychotics work? They block dopamine receptors, especially the D\(_2\) receptor subtype; this prevents endogenous dopamine from producing its normal effects.

III. Antidepressant drugs: used to treat depressive illnesses
   Most common:
   1. Monoamine oxidase inhibitors (MAOIs)
   2. Tricyclics antidepressants (ex., imipramine - Tofranil™)
   3. Serotonin-specific reuptake inhibitors (SSRIs ex., fluoxetine - Prozac™)

How do antidepressants work? MAOIs and tricyclics/SSRIs work through different mechanisms:
   - MAOIs block the breakdown of monoamines neurotransmitters (especially serotonin and noradrenaline); this therefore increases these neurotransmitters at synapses.
- Tricyclic antidepressants and SSRIs block the reuptake of monoamines (especially serotonin and noradrenaline – with the SSRIs being more specific for serotonin); also increases monoamines at synapses.
- Overall then, all antidepressant drugs increase the amount of monoamine neurotransmitters in the synapses.

**IV. Opiates (analgesics):** clinically used in the treatment of pain – high potential for addiction

Most common:
1. Opium poppy derivatives (ex., morphine, codeine)
2. Derivatives of morphine (ex., heroin)
3. Endogenous morphine (endorphins) = made in the body (ex., enkephalins, dynorphin, β-endorphin)

**How do opiates work?** All work by binding opiate receptors in the nervous system (mu, delta, and kappa subtypes of opiate receptors); the mu receptors appear to be especially important for reducing pain transmission in the nervous system.

**V. Stimulants:** in general, increase the activity of the nervous system

Several classes of stimulants:
1. Behavioral stimulants (ex., cocaine, “crack”, amphetamine)
2. Convulsant stimulants (ex., pentalenetetrazol, bicuculline)
3. General stimulants (ex., caffeine)
4. Psychedelics (ex., lysergic acid diethylamide [LSD], mescaline, marijuana, ecstasy, psilocybin [from some mushrooms])

**How do stimulants work?** Each subtype has different mechanism of action:
- Behavioral stimulants generally work by blocking dopamine reuptake at the synapse and some even produce dopamine release (amphetamine); increases dopamine at synapses;
- Convulsants block GABA receptors;
- A general stimulant such as caffeine works by blocking one subtype of adenosine receptor;
- Psychedelic drugs have various mechanisms of action which involve the serotonin system (LSD, ecstasy, mescaline, psilocybin) or other specific neurotransmitter receptors (the cannabinoid receptor for marijuana).

**What is substance abuse?** A pattern of drug use in which chronic and excessive intake takes a central place in one's life.

**What is drug dependence?** An advanced state of abuse in which physical and psychological dependence have developed.

**What is drug tolerance?** Decreased susceptibility to a drug that develops as a result of repeated exposure to the drug; mostly mediated by compensatory mechanisms that oppose the initial drug effect (ex., if drug causes hypothermia, opposing mechanism of hyperthermia is produced)

**What is cross-tolerance?** Tolerance that develops to one drug carries over to other drugs (often of the same family) suggesting that they act through the same target (mechanism)
What are the 3 different types of tolerance that we discussed in class?
1. Metabolic tolerance: reduced sensitivity to a drug that results from the increased ability of the body to metabolize the drug (ex., up-regulation of liver enzymes)
2. Cellular tolerance: a change that takes place in nerve cells in which the activity of neurons adjust to the excitatory or inhibitory effects of a drug (ex., receptor downregulation)
3. Learned tolerance: behavior change acquired through associative learning.

What is sensitization? Increased behavioral response to the same dose of a drug - for example, the behavioral activity of animals in response to repeated injections of amphetamine increases over time.

What is physical dependence? State in which discontinuation of drug taking will induce withdrawal syndrome (illness).

What is the withdrawal syndrome? Illness induced by the elimination or absence from the body of a drug on which a person is physically dependent.

What is psychological dependence? The most important factor in addictive behavior - includes several cues associated (conditioned) with pleasurable effects of drugs that are difficult to eliminate; mostly responsible for “compulsive” drug taking behaviors.

What is the role of learning in addiction? Involved in the development of psychological dependence.
1. Associations of places and cues with drug taking produces both conditioned drug tolerance and conditioned withdrawal effects (Siegel).
   Example: Rats that are injected with morphine repeatedly in context A will develop conditioned tolerance and withdrawal only in context A, not in their homecage. Thus, if they get injected with a high dose of morphine in their homecage, they are much more vulnerable to die from an overdose, than if they received the injection in context A.

What is reinforcement? Process by which there is an increase in the likelihood of a behavior immediately preceding the reinforcement (positive or negative). Reinforcement is most effective upon behaviors that have just happened - it’s effects wane very quickly with time (example of rats preferring a small reward immediately rather than a much larger reward, with a delay).

Who discovered reinforcement? In a classic study, Olds & Milner (1954) implanted electrodes in the brain of rodents that were aimed for the reticular formation, but accidentally implanted some animals in the medial forebrain bundle (MFB). They observed that the rats with MFB electrodes would do everything to get more stimulation. They argued that the MFB was a region involved in pleasure and reinforcement. (rats would die of starvation in order to obtain electrical stimulation of the MFB)

What are the neural systems of reinforcement? Several additional studies have shown that neurons from the ventral tegmental area, which mainly contain dopamine as a neurotransmitter, project their axons through the MFB to the nucleus accumbens, where dopamine is released.
   - If the effect of dopamine is blocked by injecting dopamine receptor antagonists in the nucleus accumbens, electrical self-stimulation of the MFB and drug taking behavior can be blocked in rodents.
What is the role of reinforcement in addiction? Most addictive drugs studied so far have strong reinforcing properties. Thus, the behaviors associated with drug taking become more likely (reinforced). This occurs through:

1. **Positive reinforcement**: Pleasurable effects of drugs that can be produced, but not always, consciously (remember the example of the detoxicated heroin addicts and their willingness to work hard for morphine injections that they consciously reported as “worthless”). Increases the likelihood of behavior just preceding reinforcement.

2. **Negative reinforcement**: Increase in the likelihood of a response that reduces or removes an aversive stimulus (in the case of addiction, attempts to stop withdrawal syndrome by injecting drug).

What is the theory of incentive sensitization? Why do humans and animals keep injecting drugs even if the drugs are no longer pleasurable? The incentive sensitization theory (Robinson & Berridge) suggests that the stimuli associated with drug taking (conditioned stimuli) become “attached” with motivational properties, which become “sensitized” (stronger) with repeated drug taking. So even if drugs lose their direct “pleasurable” qualities, the conditioned incentive stimuli “replace” the primary drug effects, being themselves very pleasurable.

- As a matter of fact, dopamine release in the nucleus accumbens is triggered by situational cues (in addicted rats) more than by the drug itself in addicted animals.

Addiction believed to develop in 3 stages:

1. Seeking the sensation of pleasure from drug taking;
2. Pleasure is linked to mental representations (cues) associated with drug taking;
3. Cues associated with drug taking cues become incentives, through sensitization process.

How are drug dependence problems treated? In general, the physical dependence and withdrawal symptoms that accompany discontinuation of drug taking can be monitored and slowly reduced by systematically reducing drug taking in a clinical setting (clinical drug rehabilitation programs - example of nicotine patches in individuals trying to stop smoking). At most, a few months are normally sufficient to stop drug taking and avoid serious withdrawal symptoms. Unfortunately, the psychological dependence is much more difficult to deal with, and often, individuals who have suffered drug dependence and find themselves in situations (cues) previously associated with drug taking will relapse into their drug problem.

Treatments of commonly abused substances:

**General adverse consequences of abused drugs:**
- Expensive habit (associated with crime);
- IV injections - health risks (HIV, hepatitis);
- Source can be doubtful and plain dangerous;
- Crosses placental barrier, making fetus depend;
- Overdose can produce death;
- Some drugs can lead to neurological disorders and brain damage.

**Common withdrawal symptoms:** from 2-3 hours until 6-7 days after last ingestion/dose;
- some withdrawal symptoms: increased fidgeting, sweat, sleep, shiver, nausea, vomiting, diarrhea, cramps, tremors, muscle spasms (legs - going “cold turkey” and “kicking” the habit!)
**Specific Treatments:**
- in some cases, drug specific treatments are available (ex. methadone maintenance for opiates);
- in general **NO GOOD TREATMENTS** for any addictive drugs!

**Hormones and Behavior**

**What is the neuroendocrine system?** It is the sum of the glands, hormones, and target tissues/organs involved in the control of bodily functions (including behavior).

**What are glands?** They are specific cell masses throughout the body that produce and secrete a variety of hormones (chemicals).

There are two types of glands:
1. **Exocrine glands**: they secrete their chemicals into ducts, which are carried to the surface of the body (ex. sweat and tear [lacrimal] glands).
2. **Endocrine glands**: ductless glands that secrete “hormones” into the general circulation (ex. pituitary and gonadal glands).

**What are hormones?** Chemicals released by the endocrine glands into the general circulation. Hormones can be of several general types:
   I. Amino acid derivatives (ex. adrenaline, produced from tyrosine);
   II. Short peptides and proteins (adrenocorticotropic hormone, ACTH);
   III. Steroids (ex. cortisol, estradiol; synthesized from cholesterol - remember, these are lipid soluble and freely cross the blood-brain-barrier and easily enters/exits all cells, including neurons).

**What are some examples of endocrine glands and their hormones?**
- pineal gland: melatonin;
- hypothalamus: vasopressin, oxytocin, prolactin;
- pituitary gland: luteinizing hormone, adrenocorticotropic hormone, follicle stimulating hormone;
- thyroid gland: thyroxin hormone;
- thymus gland: lymphokines (involved in immune responses);
- adrenal glands: cortisol;
- pancreas: insulin, glucagon;
- ovaries: estradiol, testosterone;
- testes: testosterone, estradiol.

**What is the hierarchical control of hormone release?**
The brain (hypothalamus) ultimately controls many of the hormones found in the body. This is usually regulated through “multi-step” signaling mechanisms (pituitary gland) all the way to the various glands in the body that synthesize hormones. In turn, many hormones reach back to the brain and influence various cognitive and behavioral functions.

**How is the Anterior pituitary controlled?**
Neurons of several hypothalamic nuclei produce and release hormones (“releasing hormones”) from their axons in the median eminence; the median eminence is highly vascularised by the hypophyseal artery, which transport the released hormones into the anterior
pituitary via portal veins; anterior pituitary cells respond to hypothalamic hormones by producing and releasing their hormones into the hypophyseal vein, which brings hormones into the general blood circulation (goes everywhere in the body).
- example: gonadotropin-releasing hormone from the hypothalamus produces the release of follicle-stimulating hormone and luteinizing hormones from the anterior pituitary, which target the gonads.

**How is the Posterior pituitary controlled?**

Neurons from different hypothalamic nuclei also control the posterior pituitary gland; hormone containing neurons of the hypothalamus release their hormones directly in the posterior pituitary gland; the released hormones then enter the hypophyseal vein and reach the general blood circulation.
- example: oxytocin is released from the hypothalamus (supraoptic nucleus) into the posterior pituitary, and is involved in uterine contraction during child birth and milk ejection during suckling.

**As an example of endocrine regulation, how are the gonads controlled?**

In the case of the gonads, neurons of different hypothalamic nuclei produce and release gonadotropin-releasing hormone (GnRH) in the anterior pituitary gland; GnRH induces the production and release follicle-stimulating hormone (FSH) and luteinizing hormone (LH) from the anterior pituitary cells, which reach the general circulation; FSH and LH eventually reach the gonads, inducing the production and release of estrogens, progesterone, and androgens (testosterone) from cells in the gonads (ovaries and testes).

**What are homeostatic mechanisms?** Mechanisms that keep some body functions within a narrow, fixed range.

**What are the 3 main mechanisms involved in homeostatic regulation?**

1. **Negative feedback:** A signal (usually a hormone in the endocrine system) in one direction that results in a compensatory effect in the other direction. Practically, when a hormone is released, it goes back to several body areas (pituitary, hypothalamus) involved in its release to terminate further release.
   - recall analogy with heating system at home, with the thermostat controlling the temperature, and high temperature shutting down the furnace.

2. **Neural control:** Neurons from other brain regions send their axons to hypothalamic nuclei and can regulate the activity of hypothalamic releasing-hormone neurons;
   - this can increase or decrease hormone release and overall levels in bloodstream.

3. **Experience/learning:** Repeated experience (learning) with a specific situation can lead to increased or decreased hormone levels;
   - often regulated by brain regions that send axons to hypothalamus.

**Motivated behaviors are often broken down into regulatory and non-regulatory behaviors.**

**What are regulatory behaviors?** Behaviors controlled by a homeostatic mechanism
Examples are temperature regulation, eating and drinking, salt appetite, waste elimination, etc.
**What are non-regulatory behaviors?** Behaviors that are not controlled by homeostatic mechanisms - all behaviors excluding those regulated by homeostatic processes. Examples are sexual behavior, parental behaviors, aggression, playing sports, etc. Both regulatory and non-regulatory behaviors are controlled by the hypothalamus.

**Example of non-regulatory behaviors - Sex!**

**What are gonadal hormones?** They include two main classes of steroid hormones (derived from the fat molecule cholesterol) known as androgens and estrogens, and a third class known as the progestins (the most common of which is progesterone); these hormones have organizational and activational effects.

**What are androgens?** The class of steroid hormones that includes testosterone.

**What are estrogens?** The class of steroid hormones that includes estradiol.

**Organizational effects of sex steroids on the brain and body:**
Process whereby gonadal hormones act on the brain during development to produce distinctly female or male brains and bodies; some brain structures differ between males and females; these structures are referred to as sexually dimorphic.
- the suprachiasmatic nucleus and parts of the preoptic nuclei (medial nucleus) are generally larger in males.
- this is produced by the organizational effects of testosterone during fetal development (that is, if testosterone is present, facilitates the development of the “male” brain); testosterone enters the nervous system and neurons of the brain and is converted to estradiol by aromatase enzyme, and leads to the development of male brain characteristics. Higher levels of estradiol in females cannot enter neurons because they are tightly bound to the protein alpha fetoprotein.
During adolescence, sex steroids rise and lead to the male and female body characteristic differences.

**Activational effects of sex steroids:**
Cycling levels of sex hormones in females lead to menstrual cycle and changes in brain and behavior throughout the cycle (remember receptivity and dendritic spines in hippocampus) whereas in males, high levels of testosterone are associated with increased motivation to seek sexual behaviors (and aggression!!!).

**What are the main brain regions associated with sexual behaviors (remember this is mostly for rodent behavior)?**
1. The ventromedial hypothalamic nucleus controls the sexual receptivity posture (mating posture - lordosis) in females;
2. The preoptic area of the hypothalamus controls copulatory behavior (thrusting) in males, but not sexual motivation;
3. The medial nucleus of the amygdala in males (and possibly females) controls sexual motivation (remember the ceiling trap door experiment!!!).
Example of regulatory behaviors - Eating and Drinking

**How is eating (feeding) controlled?** Eating is partly controlled by the digestive tract, the hypothalamus, and cognitive factors.

**What is the role of the digestive tract in the control of eating?** It functions as a reservoir where a variety of chemicals and enzymes released (ex., hydrochloric acid, pepsin, etc.) help breakdown and absorb essential nutrients and energy molecules from food.

**What are the 3 classes of molecules ingested by our body?**
1. Lipids (fats - provides majority of energy stores - 85%)
2. Amino acids (essential building blocks of proteins - provides about 14% of energy)
3. Glycogens (starches and sugars - glucose - provides about 1% of energy stored)

**What are the brain areas involved in the long-term control of feeding?** There are 3 main hypothalamic nuclei critically involved in the control of feeding:
1. Paraventricular nucleus: involved in the release of ACTH and thyrotropin from the anterior pituitary (mostly involved in metabolic regulation)
2. Lateral hypothalamus: coordinates motor responses involved in feeding behaviors
3. Arcuate nucleus: receives peripheral signals (leptin and insulin) about the state of the energy stores in the body, and projects to paraventricular nucleus and lateral hypothalamus to increase or decrease feeding

**What are the mechanisms involved in the short-term regulation of feeding?** There are signals for satiety and for hunger:

**I. Hunger signals:**
- Sight of food and thinking about food: raises activity of autonomic nervous system, which activates pancreas to release insulin in blood circulation - produces a drop in blood glucose that initiates a response in the nucleus of the solitary tract (see below)

- Hypoglycemia: a drop in blood glucose levels that is sensed by specialized receptors onto neurons of the area postrema (no blood-brain-barrier), and peripherally in the liver, via the vagus nerve (neurons in the nucleus of the solitary tract in the medulla)

- Lipoprivation: a drop in fatty acid levels available to cells, detected in liver and other abdominal organs. Information provided to brain via vagus nerve

**II. Satiety signals:**
- Gastric distension: somatosensory receptors located in gastrointestinal tract and enteric nervous system sense stomach distension, and provide this information to nucleus of the solitary tract via the vagus nerve

- Cholecystokinin (CCK): peptide released from gastrointestinal tract and enteric nervous system - responds to both volume and caloric content of food being absorbed - acts on vagal somatosensory receptors and synergises with gastric distension signals from gastrointestinal tract
- High levels of insulin (released from b-cells of pancreas) and glucose in blood serve as a satiety signal at the level of the hypothalamus (arcuate nucleus)

**With regard to water balance, how much of the human body is water?** Approximately 70%.

**What are the two mechanisms responsible for water balance (homeostasis)?**

1. **Osmotic thirst (cellular dehydration):** Increase in concentration of dissolved substances in the blood (hypertonicity)
   - Specialized hypothalamic neurons in area called vascular organ of the lamina terminalis (OVLT) sense hypertonicity
   - OVLT neurons send their axons to paraventricular nucleus neurons that produce vasopressin (antidiuretic hormone – ADH) that, when activated, release vasopressin in posterior pituitary

2. **Hypovolemic thirst:** Decrease in overall blood volume
   - Increases water retention via two mechanisms:
     a) Reduced blood flow in renal system produces release of angiotensin II from kidneys which is sensed by hypothalamic subfornical organ.
        - Subfornical organ activates paraventricular ADH neurons to release vasopressin from posterior pituitary.
     b) Mechanoreceptors (baroreceptors) in wall of large blood vessels and heart signal loss of blood to the hypothalamus via vagus nerve and nucleus of the solitary tract (NTS).
        - NTS also projects to paraventricular nucleus and synergizes with effects of angiotensin and activate sympathetic nervous system to help correct the reduction in blood pressure (produces vasoconstriction)
        - NTS also projects to lateral hypothalamus to stimulate drinking behaviors (finding water, ingesting fluids).

**Emotions**

**What are the early theories of emotions?**

I. **James-Lange theory (1884):** A theory of emotion suggesting that behaviors and physiological responses are directly elicited by situations and that feelings of emotions are produced by feedback from these behaviors and responses.
   - Example: sight of a bear elicits increases in autonomic (heart rate, blood pressure, etc.), and behavioral responses (running), which in turn produce the conscious feeling of fear.

II. **Cannon-Bard theory (1900’s):** A theory suggesting that emotional experiences and emotional expression (autonomic, hormonal, and behavioral responses) occur in parallel and have no direct causal relation to one another.
   - Example: sight of a bear elicits, at the same time, the feeling of fear and the autonomic, hormonal and behavioral responses.
What are the predictions of these two theories?

The James-Lange theory predicts that without autonomic, hormonal, and somatic feedback, a person would not feel emotions.

The Cannon-Bard theory predicts just the opposite, that person does not need feedback to feel emotions.

Studies of patients with various spinal cord injuries have lent support to both theories, while pointing to their limitations.
- patients with injuries which remove most feedback from autonomic and somatic origin still experience emotional “feelings”;
- however, these patients report much different emotional experiences, being weaker, of diminished intensity.

What is Damasio’s somatic marker theory of emotions?

Suggestion that signals (markers) arising from internal and external environment (emotional stimuli) act to guide behavior and decision making, usually in an unconscious process.

What is the brain system associated with emotions?

An early influential theory by Papez (1937) defined the “limbic system” as the brain circuit involved in the elicitation of emotional expression and feelings.
- The limbic system consists of several interconnected structures, the most important of which include the hippocampus, amygdala, cingulate cortex, fornix, septum, mammillary body, and some hypothalamic nuclei.
- Papez proposed that emotional expression was produced by limbic activity upon the hypothalamus, whereas emotional experience (feeling) was produced by limbic activity upon the cortex.
- Papez was right, to some extent, although the concept of the limbic system has greatly evolved since his first proposal; the role of the amygdala in emotions has particularly received a lot of attention.

What is the Kluever-Bucy syndrome?

It is a cluster of behaviors originally observed in monkeys sustaining bilateral anterior temporal lobectomies (also observed with amygdala lesions, and later observed in humans).

The behaviors included:
- flat affect (emotions);
- consumption of nearly everything edible;
- increased, but inappropriate, sexual activity (often towards inanimate objects);
- tendency to repeatedly investigate same objects;
- tendency to investigate objects with mouth;
- increased tameness, lack of fear.

What are fear reactions?

Fear reactions are characterized by a constellation of autonomic, hormonal, and behavioral responses preparing an organism for fighting or fleeing.

What is the brain area that regulates this constellation of reactions?

The central nucleus of the amygdala, which has connections to most brain areas controlling specific aspects of the fear reactions.
What has been learned about fear in humans?

Patients evaluated for neurosurgery and subjected to electrical brain stimulations show autonomic signs of fear when stimulated in the hypothalamus; however, they do not report any fearful “experiences”; however, amygdala stimulation elicits both autonomic and experiential fear reactions.

- patients with bilateral amygdala damage respond to fear provoking stimuli in a much blunted way; they also cannot use negatively charged emotions to remember details of a fearful situation.
- brain imaging experiments in human subjects shown frightening movies or slides show particularly high activity levels in the amygdala.

How was stress originally described?

Hans Seyle (1940’s) described a set of physiological effects related to a “generalized” alarm system in response to illness and surgery.

How is stress now defined?

A physiological reaction caused by the “perception” or “detection” of aversive or threatening situations that jeopardize some homeostatic function.

What are some examples of stressful situations?

Stressors normally fall into two classes:

1. Psychological stressors:
   - marriage, divorce, job loss, new job;
   - entering college;
   - death of a loved one;
   - public speech;

2. Systemic stressors:
   - limb fracture;
   - surgical procedures;
   - viral or bacterial infections;
   - food poisoning;
   - illnesses (cancer, heart conditions, etc.).

What are the physiological responses to stressful situations?

There are two different sets of responses; one produced by acute stressors (normally advantageous), and one produced by chronic stressors (normally disadvantageous):

1. A cute stress responses:
   - activation of autonomic and endocrine responses;
   - release of adrenaline and noradrenaline by the adrenal medulla;
   - release of cortisol by the adrenal cortex;
   - increases in heart rate and blood pressure to bring more blood to muscles;
   - mobilization of energy resources (production and release of glucose for use by muscles;
   - inhibition of inflammatory responses;
   - resistance to infection;
   - inhibits sexual functions and sex steroid production and secretion.

2. Chronic stress responses:
   - hypertension (high blood pressure);
   - gastrointestinal ulcers;
- diabetes;
- inhibition of growth (particularly important in young children – psychological dwarfism);
- infertility;
- suppression of the immune system;
- damage to the brain (hippocampus).

How does the brain control the autonomic and endocrine responses to stress?

Sympathetic neurons of the autonomic nervous system located in the brainstem innervate and activate the secretory cells of the adrenal medulla to secrete adrenaline and noradrenaline in the general circulation.

The hypothalamus (paraventricular nucleus) releases corticotropin-releasing hormone (CRH) in the median eminence, which is transported to the anterior pituitary and elicits the release of adrenocorticotrophic hormone (ACTH) in the general circulation, which produces the release of glucocorticoids (cortisol) in the general circulation from the adrenal cortex.

What brain areas are involved in the perception of different stressful situations?

1. Limbic system: involved in the perception of “psychological stressors”;
2. Circumventricular organs: involved in the detection of “blood born pathogens” – poisons;

All these areas converge upon the paraventricular nucleus of the hypothalamus, which controls the anterior pituitary via the release of CRH and the brainstem autonomic nuclei by direct neural connections.

How can one control stress?

A lot of evidence suggests that the perception of control over stressful situations reduces several physiological reactions produced by these stressful situations. The perception of control produced by behavioral or mental responses is termed the coping response.