

Pain Pathways Ma	ade Sim	ple
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Disclosures

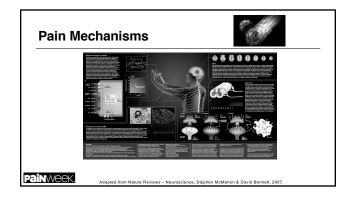
Nothing to Disclose

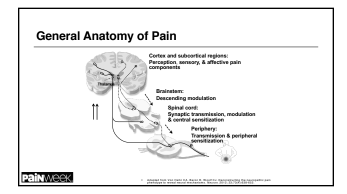
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Learning Objectives

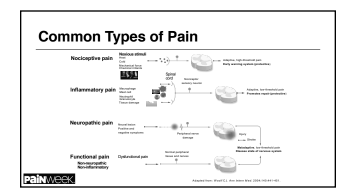
- Differentiate between nociceptive and neuropathic pain
- Describe the process of pain transmission
- Identify the specific pain pathways that can be acted upon by common pharmacotherapy classes

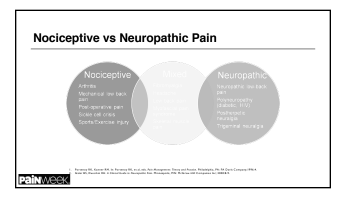
Classification of Pain	
■Good pain vs bad pain	
>	
Clinical Pearl	
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Good Pain	
Nociceptive pain: purposeful pain	
 -Eudynia — pain linked to normal tissue function or damage -Nonmaldynic pain 	
-Adaptive	
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Bad Pain	
Neuropathic pain: nonpurposeful pain	
-Neuropatific pairt frompurposerul pairt -Maldynia—pain linked to disorder, illness or damage	
-ie, may be abnormal, unfamiliar pain, assumed to be caused by	-
dysfunction in PNS or CNS	
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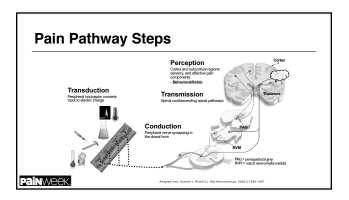




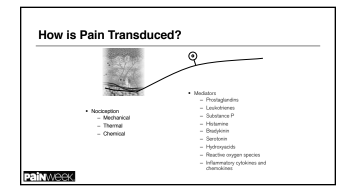
Pain Roadmap: Peripheral and Central Nervous System Landmarks Physiologic process involving multiple areas of the nervous system Bidirectional Involves normal as well as pathological processes A sensory experience associated with affective and cognitive responses Dynamic (ie, occurring in real time) Adapts or changes in response to function—"neuroplasticity"

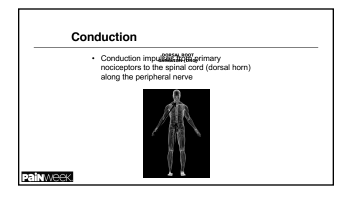






Transduction: Processing at Peripheral Nerve Endings	
Conversion of mechanical, thermal or chemical stimuli into an electric charge Involves Receptors activated directly by stimuli Injury/inflammatory response	
Adapted from Disciplanty PM, et al. Neuroschemistry and pain processing in: Binezzo N, et al., et al., et al. Essentials of Pinis Medicine. Philodophia, PA. Saundere, 2011. chapter 2	





Primary Nociception

- A-delta fibers
 - -Small receptive fields
 - Thermal & mechanical
 - Myelinated
 - Rapidly conducting
 - 10-30 m/sec
 - Large diameter



C-fibers

- Broad receptive fields
- Polymodal
- Unmyelinated
- Slower conducting
- .5-2.0 m/sec - Cross sensitized
- -Small diameter



Peripheral Pain Nociceptors

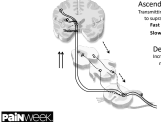


 $A\beta-$ muscle spindle secondary endings, touch, and kinesthesia $A\delta-$ pain, temperature, crude touch, and pressure

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Bashbasum A, Jessell T, The perception of Pain, In Kendal E, Schwartz J, Principles of Neural Science 4° ed, New York, IntGraw Hill, 2000, 482-483.

Transmission & Modulation



Ascending nociceptive pathways
Transmitting nociceptive impulses from the dorsal horn smitting nociceptive impulses from the dorso o supraspinal targets Fast (green) neospinalthalamic Slow (yellow) paleospinalthalamic

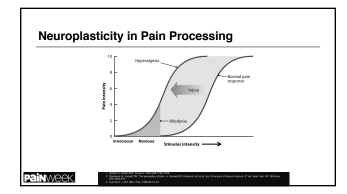
Descending inhibitory tracts (blue) Increase or decrease volume control of incoming nociceptive signals reaching the brain 5-HT—serotonin

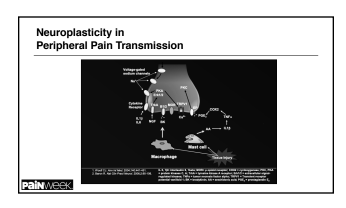
NE—norepinephrine

Adapted from Von Hehn CA, Baron R, Woolf CJ. Deconstructing the neuropathic pain phenotype to reveal neural mechanisms. *Neuron*. 2012; 23;73(4):638-652.

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Harris Bain Conducted and Transmitted	
How is Pain Conducted and Transmitted?	
ORSAL ROOT IGLION	
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• Inhibitory transmitters	
- Substance P (descending inhibitory pathways) - GABA	
- Calcitonin gene related peptide - Glycine - Aspartate, glutamate - Somatostatin	
— α2 agonists	
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Role of Neuronal Plasticity in Pain	
 Nervous system changes in Neuronal structure 	-
- Connections between neurons	
 Quantity/properties of neurotransmitters, receptors, ion channels Decreases body's pain inhibitory systems (increased pain) 	
 Injury, inflammation, and disease are culprits Produces short-term and permanent changes 	
Pivotal to the development of hypersensitivity of inflammatory pain	
■ Enables NS to modify its function according to different conditions or demands	
placed upon it	
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How Acute Pain Becomes Chronic	
■ Peripheral sensitization	
 Tissue damage releases sensitizing "soup" of cytokines & 	
neurotransmitters	
COX-mediated PGE2 release Sensitized nociceptors exhibiting a decreased threshold for activation &	
increased rate of firing	
Central sensitization—resulting from noxious input to the spinal cord Resulting in hyperplaceia & alledunia	
Resulting in hyperalgesia, & allodynia	

■ Hyperalgesia - Lowered threshold to different types of noxious stimuli ■ Allodynia - Painful response to what should normally be nonpainful stimuli





Periphe	eral Sensitization	
	Sensitizing agent Mechanical Chemical Thermal PLC Thermal Thermal Thermal Thermal Ion channel phosphorylation Generator potential (membrane depolarization) Thermal Freech voltage sensitive channel threshold	
	Action potential	
	Adapted from: Griffin RS, Woolf CJ. The Pathophysiologic Basis of Drug Therapy, In: Golan DE, Tashijan AH, Armstrong EJ, Armstrong AW (eds), Principles of Pharmscodigy, Baltimore, MD, Lippincott Williams & Wilkims, 2005;235-246.	
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Central Sensitization

- Activation
 "Wind up" of dorsal horn nociceptors
- Modulation
 Excitatory/Inhibitory neurotransmitters
- Decreased central inhibition of pain transmission

Prime role in chronic pain, particularly neuropathic pain

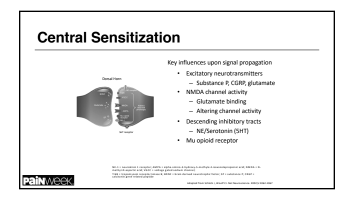
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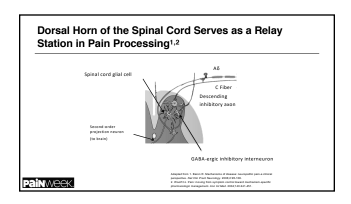
Definitions

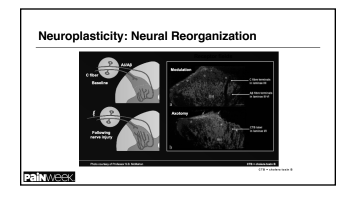
- Wind Up
 - Causes long-term changes in nociceptive neurons, which become hyperexcitable such that they respond to lower stimuli
 - NMDA-type glutamate receptors play an important role in this process^{1,2,3,4}
 - Prolonged opening of the ion channels enables greater influx of calcium and sodium across the post-synaptic membrane and greater excitation of nociceptive neurons^{2,3}

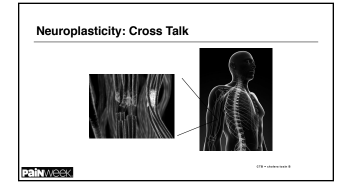
1.	Kandel ER	, Schwartz JH, Jessell TM, editors. Principles of Neural Science (Fourth Ed
		New York: McGraw Hill (Health Professions Division). 2000;472-491.

Central	Sensitization	
	Afferent first order neuron	Dorsal horn neuron
	CARAMAN SP	VOC. Adhed correct of the correct of
	NK-1 = neurokinin 1 receptor; AMPA = alpha-ami methyl-0-aspartic acid; VGCC = voltage gated sod TrkB = tropomyosin receptor kinase B; BONF = br	to-3-hydroey-6-methyle 4-lessassleproplonic acid; NM.DA + N- lum channel; als derived neurotrophic factor; SP = substance P
Painweek.		Adapted from Schlotz J, Woolf CJ. Nat Neuroscience. 2002;5:1063-1067







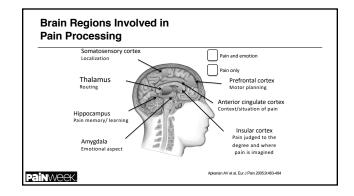


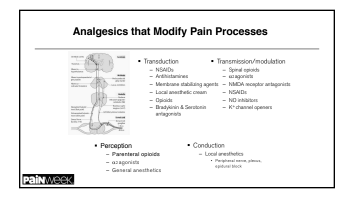
Central Sensitization: Neuroplasticity in Spinal Cord Processing • Definition: altered function of neurons or synaptic activity • Mechanisms of central sensitization may include: - Changes affecting glutamate/NMDA receptors activity • Reduced threshold for activation • Increased availability of glutamate • Increased influx of Nar/Car (receptor open longer) - Modulation—excitatory/Inhibitory neurotransmitters - Decreased tone—descending inhibitory pathways² - Activation/migration of glial cells into the spinal cord³

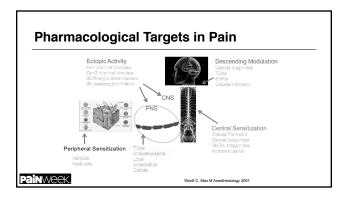
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Mannion R.J. Woolf C.J.: Clin J Pain. 2000;16(3):S151-S153. 2. Oxsigov MN, et al. Ann NY Acad Sci. 2000;900:12-24
 Waxeler-Frank J, et al. Neurosignair. 2005;14:166-174. 4. Guilbaud G, et al. Exp Brain Rez. 1992;92:227-243.

- Changes in the thalamus and primary somatosensory cortex⁴





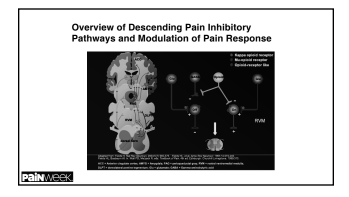


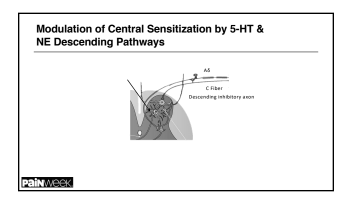
The Chronic Pain Armamentarium Nonopiolds - Acetaminophen - NSAIDs - COX-2 inhibitors Opiolds - Mu-opioid agonists - Mixed agonist-antagonists Adjuvant analogsics - Antidepressants - Anticonvulsants - Topical agents/local anesthetics

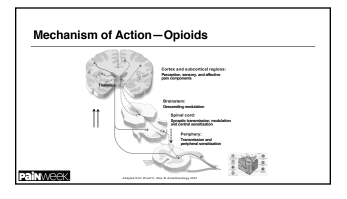
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Common Pharmacologic Therapies - Acetaminophen - NSAIDS - Antiepileptics - TCAs - SNRIs - Topicals - Muscle relaxants - Opioids

Nonopioids: Acetaminophen	
Example	
- Acetaminophen	
Mechanism of action	
-Inhibits prostaglandin production in CNS; antipyretic activity	
 No effect on blocking peripheral prostaglandin production; no anti-inflammatory or antirheumatic activity 	-
FDA warning	
Potential severe liver damage if over-used	
 Stevens-Johnson Syndrome & toxic epidermal necrolysis 	
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Nonopioids: NSAIDs	
<u>Examples</u> - Acetylated (aspirin); nonacetylated (diflunisal);	
acetic acid (diclofenac); propionic acid (naproxen); fenamic acid (mefenamic acid);	
<pre>(naproxen); fenamic acid (mefenamic acid); enolic acids (piroxicam); nonacidic (nabumetone);</pre>	
ibuprofen, selective COX-2s (celecoxib)	
Mechanism of action	
 Exhibit both peripheral and central effects; anti-inflammatory and analgesic effects 	
 Inhibition of cyclooxygenase and prostaglandin 	
production Inhibition of leukotriene B4 production	
- Lipoxins (signaling resolution of inflammation)	
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Opioids	
<u>Examples</u>	
-Morphine, hydromorphone, fentanyl, oxycodone, oxymorphone,	
meperidine, codeine, methadone, tramadol	
Mechanism of action - Bind to opioid receptors in the central nervous system (CNS) to	
inhibit transmission of nociceptive input from periphery to spinal cord	
 Activate descending pathways that modulate transmission in 	
spinal cord - Alter limbic system activity; modify sensory and affective	
pain aspects	







Adjuvant Analgesics: Tricyclic Antidepressants

Examples

- Amitriptyline, desipramine, doxepin, imipramine, nortriptyline Mechanism of action
 - Reduction in action potential firing of sodium channel activity
 - Inhibition of reuptake of NE and 5-HT
 - Analgesia is independent of antidepressant function
 - High side effect profile (tolerability)
 - Cardiotoxic (overdose)

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TCAs and SNRIs Pharmacological Properties | Topic reference backet | T

SSRIs (Selective Serotonin Reuptake Inhibitors)

Examples

-Citalopram, fluoxetine, fluvoxamine, paroxetine, and sertraline Mechanism of action

-Selectively inhibit 5-HT reuptake without affecting NE

Therefore, no pain relief expected!

Serotonin	
 International Union of Pure and Applied Chemistry nomenclature 5-Hydroxytryptamine (5-HT) 	
- Monoamine neurotransmitter, biochemically derived from tryptophan	
 Receptors are a group of G protein-coupled receptors (GPCRs) and ligand-gated ion channels (LGICs) found in the central and peripheral 	
nervous systems	
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Serotonin/5-HT Receptors	
Family Type Mechanism Potential	
S-HT (Q/G_presis coupled Decreasing cultural rovels of CAMP. Including Cultural rovels of CAMP. S-HT (Q/G_presis coupled Increasing cultural rovels of FD and DAG. Existatey S-HT (Ligard-pared Na" and N" crisin channel. Operbrining plasma membrane. Excitatey	
SHTa Gu-protein coupled Increasing offshire levels of cAMP Excitatory SHTB Q/Gu-protein coupled The Camp Increasing offshire levels of CAMP Excitatory SHTB Gu-protein coupled Increasing collabal revised CAMP Excitatory SHTB Gu-protein coupled Increasing collabal revised CAMP Excitatory	
SHI Op-protein coupled. Increasing centure revers of centure Landautry	
PainWeek. http://en.wikipedia.org/wiki/5-HT_receptor	
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Serotonin/5-HT Receptors	
5-HT1a (blood ves/CNS) 5-HT1a (cont'd)	
- Addiction - Respiration - Aggression - Sexual behavior	
- Anxiety - Sleep - Appeilie - Sociability - BP	
Cardiovascular function — Thermoregulation Emesis Heart rate Heart rate	
- Impulsivity - Sleep - Memory - Anxiety	
- Mood - Cognition - Nausea - Learning - Nociception - Memory	
- Penile erection - Mood - Pupil dilatation	
PainWeek. http://en.wikinedia.org/wiki/5.HT.recentor	

SNRIs (Serotonin/Noradrenaline Reuptake Inhibitors)

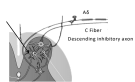
Examples

-Duloxetine, milnacipran, and venlafaxine

Mechanism of action

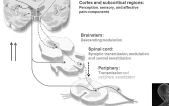
- -Block reuptake of 5-HT and NA
 - (Better tolerated, lower tendency for drug-drug interactions, better overdose safety)

Modulation of Central Sensitization by 5-HT & NE **Descending Pathways**



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Site of Action-SNRIs





Adjuvant Analgesics: Antiepileptics

Examples

 Gabapentin, pregabalin,* carbamazepine, phenytoin, divalproex sodium, clonazepam, levetiracetam, topiramate, lamotrigine

Mechanism of action

- -Suppress neuronal hyperexcitability via
 - Reducing neuronal influx of sodium (Na+) and calcium (Ca+ +)
 - Direct/indirect enhancement of GABA inhibitory effects
 - Reduce activity of glutamate and/or blocking NMDA receptors
- \bullet Binds the $\alpha2\delta$ subunit of voltage gated Ca+ channels, inhibit NT release

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Site of Action—Antiepileptics Spinal cord glial cell Spinal cell Spinal cord glial cell Spinal cel

Adjuvant Analgesics: Topicals

Examples

- $-\,\text{Lidocaine}$ patch 5% , eutectic, mixture of lidocaine and prilocaine
- Capsaicin cream/patch
- Diclofenac (cream/liquid/gel/patch)

Mechanism of action

- Block sodium channels and inhibit generation of abnormal impulses by damaged nerves
- Depletion of peripheral small fibers and therefore substance P release from sensory nerve endings
- Target local inflammatory response

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- Decrease tone of skeletal muscles
- Subclasses
 - Neuromuscular blockers
 - Act at the neuromuscular junction
 - Often used in surgery to cause temporary paralysis
 - -Spasmolytics
 - Centrally acting

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Muscle Relaxants—Spasmolytics

- Enhancing the level of inhibition
- Mimicking or enhancing the actions of endogenous inhibitory substances, such as GABA
- Reducing the level of excitation
- Common examples
 - Cyclobenzaprine (TCA) methocarbamol, carisoprodol (barbiturate like effects), tizanidine (α-2 agonist), baclofen (GABA agonist), orphenadrine (diphenhydramine/antihistamine)
- Common adverse effects
 - Sedation, lethargy & confusion (cyclobenzaprine), dependence (carisoprodol)

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Case Study





Importance for Understanding Pain Mechanisms

- Allow for rational rather than empirical approach to pain control
 Foster the development of diagnostic tools to identify specific pain mechanisms
 Facilitate pharmacotherapies that act on specific pain pathways and mechanisms
- and mechanisms

 Reduce the number of pharmacotherapies and incidence of drug-related adverse events (rationale polypharmacy)

 Enhances use of nonpharmacologic treatments

 Improve overall patient care and outcome

 Tailoing treatment based on the individual patient and pain type

 Do not forget to look for the spear

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