The physiology of pain: an update and review of clinical relevance

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Introduction

Pain, defined as a symptom, sign or a syndrome, has been researched extensively compared to any other area in neurophysiology. This short review aims to explain the neurophysiological processes involved in the perception of pain, including some of the recent findings, which have led to novel targets of pain relief.

Classification

Pain is a complex perceptual experience. It has been classified based on pathophysiology, type of stimulus, site, duration, and severity. A recent, widely accepted pathophysiological classification classifies it as nociceptive, neuropathic and psychogenic.¹,² Sensation evoked as a result of localized stimulation of pain receptors (i.e. nociceptors) via mechanical, chemical or thermal modalities is referred to as nociceptive pain. Pain occurring as a result of abnormal signaling due to an injury or dysfunction of peripheral nociceptive neurons is classified as neuropathic pain. Psychogenic pain is the occurrence of pain sensation as a result of a psychiatric disorder in the absence of injury or inflammation in the affected site.

Neuroanatomy

Irrespective of its pathophysiology or categories, pain is commonly perceived as an unpleasant sensation. It is initiated by a noxious stimulus that generates impulses, which then ascend in pathways in the spinal cord to transmit information from the nociceptors to the brain.

The nerve endings containing the nociceptors are the primary afferents, terminating in the dorsal horn of the spinal cord. These nerves are small myelinated A delta fibers and unmyelinated C fibers. Following the stimulation of nociceptors, the generated action potential is propagated along these primary afferents, resulting in the release of excitatory amino acids (e.g. glutamate and aspartate), neurotrophins (e.g. Brain Derived Neurotrophic Factor; BDNF) and peptides such as substance P (SP), Neurokinin A and Calcitonin Gene Related Peptide (CGRP) from the nerve endings in the dorsal horn of the spinal cord.³,⁴,⁵

There are multiple neuronal networks involved in sensory-discriminative component (i.e. intensity, location, duration, temporal pattern etc.) and affective-cognitive component (i.e. relationship between pain with mood, memory, tolerance etc.) of pain sensation. Recent neuro-imaging techniques have been successful in defining some of the cortical regions involved in these multiple circuits. The somatosensory area of the post-central gyrus, ventroposterolateral and ventroposteromedial nuclei of the thalamus are found to play a major role in the sensory-discriminative component of pain.⁵,⁶ The somatosensory area II in the lateral parietal cortex, the ventromedial and ventroposteriorinferior nuclei of the thalamus, the inferior parietal cortex, the insular cortex, the amygdala and hippocampus play a part in the affective-cognitive component.⁵,⁶ Furthermore, the lateral spinothalamic tract, which is the ascending pathway in the spinal cord, contains discrete pathways to carry sensory-discriminative (neospinothalamic) and cognitive-affective (paleospinothalamic) components of pain.⁵,⁶,⁷,⁸

The communication between the neurons of the pain pathways is mainly via chemical neurotransmitters. Several neurotransmitters and their receptors transmit and modulate the sensation of pain.

Chemical transmission

A painful stimulus causes inflammatory changes at the site of stimulation, resulting in activation of inflammatory mediators and immune cells. The immune cells produce cytokines and prostaglandins, which act on nociceptors to depolarize the cell membrane and generate action potentials, which can be transmitted along the nerve fibers of the pain pathway. The depolarization of the nociceptive membrane is due to the opening up of voltage and ligand-gated ion channels (e.g. Na⁺, Ca²⁺, P₂, transient receptor potential-TRP). The sensitized immune cells also secrete nerve growth factors (NGF, BDNF), which alter the sensitivity of the nociceptors by acting on tyrosine kinase receptors.³,⁴,⁹ Lowering the threshold of the
nociceptors leads to a prolonged response to stimulation. In the presence of nerve damage following injury, the degenerating nerves release CGRP and substance P, which enhance the production and release of more inflammatory mediators, thus leading to prolonged pain sensation.\textsuperscript{10} BDNF activates tyrosine kinase receptors B in the secondary afferents in the spinal cord.\textsuperscript{5} Substance P (SP) acts on Neurokinin 2 receptors in the secondary afferents and contribute to the induction of dorsal horn sensitization.\textsuperscript{5,11} CGRP causes Ca\textsuperscript{2+}-influx and prevents the metabolism of SP, and increases the release of SP and excitatory amino acids. Thus, CGRP strengthens the process of sensitization.\textsuperscript{3,4} The excitatory amino acids, especially glutamate, bind with the receptors including ionotropic receptors such as AMPA (-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid), NMDA (N-methyl-D-aspartate), Kainate and metabotropic (acting via G-proteins to soluble second messengers) glutamate receptors in the presynaptic and postsynaptic second-order neurons in the pain pathway.\textsuperscript{10,12} The NMDA receptors are considered to play an important role in the transmission of chronic persistent pain and have been found responsible for the induction and initiation of hyperalgesia (i.e. increased pain sensation to a mild noxious stimulus) and allodynia (i.e. pain sensation to an innocuous stimulus). It has also been suggested that hyperalgesia and allodynia occur due to the rewiring in the dorsal horn, with abnormal sprouting of A-beta fibers which forms synapses with A-delta and C fibres, bypassing the inhibitory interneurons and the descending modulatory pathways.\textsuperscript{13,14} Inhibitory interneurons secreting gamma amino butyric acid (GABA) modulate pain transmission at the superficial layers of the dorsal horn by presynaptic inhibition of pain fibers, thus preventing the release of excitatory amino acids. In an injury, especially with neuronal damage, the GABAergic influence is diminished due to increased release of excitatory amino acids and also hyperactivation of NMDA receptors, which can cause apoptosis of inhibitory interneurons.\textsuperscript{15}

In the presence of persistent inflammatory stimulation, changes occur in genetic regulation, which leads to induction of new proteins and modulation of the levels of expression of existing neurochemicals mediating pain.\textsuperscript{5,13} This transcription-dependent central sensitization may explain the persistent refractory pain syndromes which are sometimes independent of peripheral noxious stimulation and resistant to treatment.

### Modulatory pathways

The human nervous system is equipped with innate inhibitory and excitatory mechanisms and pathways to modulate the sensation of pain. The majority of the mechanisms are inhibitory and are activated with the initiation of nociceptive information to alleviate pain. The descending pathways originating from cortical and sub-cortical areas modulate transmission of pain, mainly at the dorsal horn of the spinal cord.

The descending noradrenergic pathway from the locus coeruleus stimulates adrenergic receptors (mainly alpha-2) in the dorsal horn, resulting in a decrease in the release of substance P, hyperpolarizing the sensory afferents and depolarizing the GABAergic neurons to inhibit pain transmission.\textsuperscript{16,17} The descending serotonergic pathway from the nucleus raphe magnus has both inhibitory effects (by activating 5-HT1A and 5-HT7 receptors) and excitatory effects (via 5-HT2A and 5-HT3 receptors) on pain transmission.\textsuperscript{18,19}

The opioid system is a major inhibitory system of nociception in the body, containing receptors for endogenous and exogenous peptides referred to as \(\beta\)-endorphins, enkephalins and the dynorphins. The receptors for these opioids are distributed in the afferents in the peripheral nerves, spinal cord and the brain. Up to seventeen classes of opioid receptors have been postulated, out of which three have long been confirmed to be functionally important, with identified genes responsible for their expression.\textsuperscript{20,21} These receptor classes are \(\mu\)-mu, \(\delta\)-delta and \(\kappa\)-kappa. Apart from the three major classes, receptors such as sigma, nociceptin and toll like receptor are also found to be involved in the analgesic effect and the side effects (e.g. tolerance and dependence) of opioids.\textsuperscript{20,21}

The opioid peptides modulate nociceptive input in two ways: 1) block neurotransmitter release by inhibiting Ca\textsuperscript{2+} influx into the presynaptic terminal and/or 2) open potassium channels, which hyperpolarizes neurons and inhibits spike activity. High densities of opiate receptors are found in periaqueductal gray (PAG), nucleus raphe magnus (NRM), and dorsal raphe (DR) in the rostral ventral medulla, caudate nucleus (CN), septal nucleus, hypothalamus, habenula, hippocampus and the dorsal horn of the spinal cord.\textsuperscript{21} Opioid synthesis and activity have been found to be increased following acupuncture and placebo induced pain relief.\textsuperscript{22,23,24}

The cannabinoid system exerts an on-demand nociceptive effect, with the receptors distributed in the CNS and periphery. The primary antinociceptive target of the endocannabinoids (i.e. AEA, arachidonylethanolamide; anandamide, 2-AG,2-arachidonylglycerol) and exogenous cannabis is the CB1 receptor found in abundance in PAG, RVM, dorsal horn and the periphery.\textsuperscript{22,23} CB1 is a G-protein coupled pre-synaptic receptor, which exerts its effects via the inhibition of neurotransmitter release by blockade of voltage dependent calcium channels and activation of potassium channels.\textsuperscript{22,23,24} Endocannabinoids are secreted by post-synaptic neurons and are metabolized
by enzymes found in the post-synaptic (e.g. FAAH, Fatty Acid Amide Hydrolase) and presynaptic (MAGL, monoacylglycerol lipase) neurons. The reuptake of endocannabinoids to the post-synaptic and uptake in to the pre-synaptic membrane for metabolism are suggested to be by a specific endocannabinoid membrane transporter, which is yet to be cloned. The CB2 receptor, which is found in the peripheral tissues, is also found to be involved in the initiation of nociceptive transmission.

New targets for pain relief

Traditional analgesics such as non-steroidal anti-inflammatory drugs (NSAIDs) and opioid agonists (e.g. morphine) are ineffective in some pain disorders and pose problems with side-effects, thus mandating search for novel therapeutic agents with superior efficacy and safety. An important target is the secretion and metabolism of neurotransmitters such as serotonin, norepinephrine, neurokinin, GABA, glutamate and their receptors such as NMDA. The ion channels such as sodium channels, N-type calcium channels, TRP and P2 receptors are increasingly being involved in studies on pain relief. The pain modulatory systems including the endocannabinoids and the CB receptors, central opioid receptors, nerve growth factors and glial cell are also potential targets for effective pain relief.

Traditional tricyclic anti-depressants (such as amitriptyline and imipramine), which inhibit the uptake of serotonin and noradrenaline from the descending analgesic pathways, are effective in neuropathic pain. However, the positive effects of the above agents may be compromised by their side-effects, due to the widespread systemic effects involving other receptors, such as the cholinergic and histaminergic, leading to cardiorespiratory effects, glaucoma and urinary retention. Newer agents such as venlafaxine and duloxetine enhance both serotonin and norepinephrine activity in the pain modulatory pathways, with less affinity for cholinergic and histaminergic receptors. Both venlafaxine and duloxetine have been found to be effective in treating painful polyneuropathies and diabetic neuropathy, with less side effects.

Anticonvulsants such as carbamazepine, sodium valproate, oxcarbazepine, topiramate, vigabatrin and levetiracetam have been found to demonstrate analgesic effects by enhancing GABA activity, inhibiting glutamate release, blocking NMDA receptors and blocking neuronal membrane Ca
dromains and Na+ channels. Carbamazepine and oxcarbazepine are effective analgesics in trigeminal neuralgia. However, the use of newer anticonvulsants has still not been proven to have superior efficacy compared to traditional analgesics in treatment of neuropathic pain or any other pain syndrome.

NMDA receptor antagonists such as dextromethorphan, amantadine, memantine and ketamine and compounds with varying degree of NMDA antagonism have been demonstrated to be effective analgesics. Such antagonists when combined with other effects such as central opioid effect (e.g. methadone), GABAergic effects (e.g., anti-convulsants) have shown varying degrees of analgesic properties and have shown high levels of efficacy in treating neuropathic pain and associated hyperalgesia and allodynia. But NMDA antagonism leads to unwanted psychological side-effects, more severe with systemic administration, which requires further research into an NMDA antagonist with high efficacy and less side effects.

Using compounds that directly target the ion channels in the peripheral and central nociceptive pathways is considered to have the potential for higher therapeutic efficacy with fewer side effects. TRPV1 antagonists such as SB-705498, AMG517, AMG628 and ABT102 have all been found to be effective in the treatment of inflammatory and neuropathic pain in animal models. Since the TRPV1 modulators affect both central and peripheral pathways, further studies would be needed to identify potential side effects (e.g. hyperthermia) prior to translational studies.

Gabapentin and a more recent analogue pregabalin are widely used as treatment for neuropathic pain. The above analgesics, earlier considered as GABA analogues, are now found to be more specifically binding with voltage-gated calcium channels and inhibiting the release of glutamate, both at presynaptic and post synaptic sites, peripherally and centrally. Combination of opioids with gabapentin has demonstrated synergistic effects in relieving neuropathic pain.

The endocannabinoid system has long been explored as an important modulatory pathway for pain relief. This system is postulated to be involved in the analgesic effects produced by some of the commonly used analgesics, including paracetamol. CB1 receptor agonists (e.g. THC, cannabidiol) have demonstrated efficacy in pain relief in chronic pain disorders. However, their use is greatly hampered by the unwanted psychotic effects. CB2 receptor agonists have recently been shown to be effective in pain relief, possibly via its peripheral stimulatory activity on immune cells. Inhibitors of the metabolizing enzymes (e.g. FAAH and MAGL inhibitors) have demonstrated efficacy in reducing pain sensitivity in animal models of acute and chronic pain. The endocannabinoid system, especially with receptor and enzyme targets in the periphery, is an attractive target for the development of future analgesics.

Activated glial cells have been demonstrated to
release pro-inflammatory mediators that contribute to neuropathic pain. Some potential targets for neuropathic pain include blocking glial cell activation, prevention of bio-synthesis of the cytokines and antagonize the action of the pro-inflammatory cytokines.\textsuperscript{2,3,1} Several animal studies have demonstrated the effectiveness of substances such as etanercept and anakinra in blocking glial cell-mediated cytokine signaling.\textsuperscript{2,3,1} The glial cells are emerging as an attractive target for treatment of chronic pain; this, however, needs a careful approach, considering the neuroprotective functions of these cells.

Adenosine triphosphate (ATP) induces pain by activating P1 and P2 receptors. The P2 receptor, which is G protein-coupled, has received attention as a potential target for the relief of acute and chronic pain.\textsuperscript{2} Compounds such as 1-benzyl-5-phenyltetrazoles and N’-acyl hydrazides have demonstrated their efficacy as antagonists to the P2 receptor sub-type P2X7.\textsuperscript{40} The P2 receptor is suggested as a very effective potential target for pain relief, with the recently developed small molecular antagonists demonstrating site-specific activity. However, further studies are needed to elucidate the side effects and adverse effects of these compounds.

**Conclusion**

The neurobiology of pain is continuously being explored and the novel therapeutic targets for pain relief are being tested. The multiple complex circuitries of the pain pathway, with central and peripheral modulation of the perception, make it a very complex sensation with sensory, cognitive and emotional components associated with it. The variety of neurochemicals and receptors involved in the perception of pain has led to the search for many targets of pain relief. Novel targets are being studied to counteract pain syndromes which are resistant to current treatment modalities. Site-specific and selective agents are receiving a lot of attention with the hope for high efficacy and minimal side effects.

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