

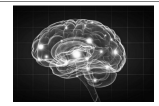
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None			

Learning Objectives

- •List the leading causes of pain after stroke. ■ Review the diagnostic criteria for central post stroke pain.
- Describe the proposed mechanisms for central post stroke pain.
 Identify a plan for medical and non-medical management for CPSP.

- Introduction
- Epidemiology
- Clinical Presentation
- Proposed Mechanisms
- Management
- Conclusion



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Central Neuropathic Pain

Common Causes:

- Ischemic/hemorrhagic stroke
- Multiple sclerosis
- Spinal cord injury
- Syringomyelia
- Vascular malformations
- Infections
- Traumatic brain injury
- Parkinson's disease?

Lancet Neurol 2009; 8: 857–68

Epidemiology

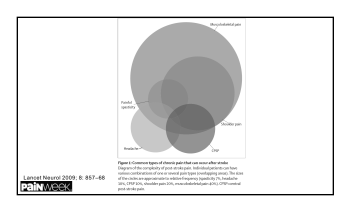
- Annually, 500, 000 people in the US have a first stroke
- ■200, 000 have a recurrent stroke
- ■80% of strokes are ischemic, either thrombotic or embolic in origin
- 5 million people in the US have had a stroke & are living in the community setting
 Of these, 1.1 million have limitations in their daily functioning or ability to perform activities of daily living
 100, 000 people have stroke as their primary diagnosis & are receiving in home health care

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- Pain is among the most common complications of stroke, with reported prevalence of 39% to 55%.
- The leading types of post-stroke pain are headaches, shoulder pain, spasticity, and central post-stroke pain (CPSP).
- Central post-stroke pain is a neuropathic pain disorder caused by the strokerelated lesion affecting the central somatosensory pathways, and accounts for about 25% of post-stroke pain cases.

PAIN 1995;61:187-93. PAIN 2011;152:818-24.

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CPSP

- ■First introduced in 1891 by Edinger.
- ■In 1906, Déjerine and Roussy provided descriptions of CPSP in 8 pts.
- •Further described by Head and Holmes in 1911 describing sensory deficits and pain narratives.
- Riddoch described symptoms of both thalamic and extra-thalamic origin (1938).



Time Course

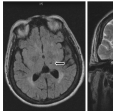
- Variable
- Can develop immediately after stroke in some patients and up to years later in others.
- Onset can be delayed, but development of CPSP within the first few months is most common.
- In a prospective study that included 16 patients with CPSP, pain onset occurred within the first month after stroke in ten patients, between 1 and 6 months in three patients, and after 6 months in three patients.
- Any later onset of pain should prompt an examination for other causes, such as a new stroke.
- Gradual onset of pain is most common. PainWeek. Lancet Neurol 2009; 8: 857–68

Diagnostic Criteria

- Mandatory criteria
- Pain within an area of the body corresponding to the lesion of the CNS.
 History suggestive of a stroke and onset of pain at or after stroke onset.
- Confirmation of a CNS lesion by imaging or negative or positive sensory signs confined to the area of the body corresponding to the lesion.

 Other causes of pain, such as nociceptive or peripheral neuropathic pain, are excluded.
- or considered highly unlikely.
- Supportive criteria
- No primary relation to movement, inflammation, or other local tissue damage.
 Descriptors such as burning, painful cold, electric shocks, aching, pressing, stinging, and pins and needles, although all pain descriptors can apply.
- -Allodynia or dysesthesia to touch or cold.

PainWeek, Lancet Neurol 2009; 8: 857-68



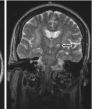


Figure 1. Axial T2 FLAIR MR image (left panel) showing a chronic left thalamic infarction (arrow). A T2 coronal image (right panel) demonstrates the postero-lateral thalamic location of the infarct.

PaiNVCCK Top Stroke Rehabil 2013;20(1):116-123.

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- Pain scales:
 - VAS or NRS are useful in the evaluation of the pain intensity, but there are no scales developed speciifcally for CPSP.
- •Quantative Sensory Testing (QST):
- -Have been used to document common or dissociated sensory findings.
 -Enable detailed sensory testing of controlled and graded physiological stimuli, such as thermal, pressure, pinprick, and vibration stimuli.

Lancet Neurol 2009; 8: 857–68

Clinical Characteristics

- Pain can be spontaneous or evoked.
- Spontaneous is common and reported in 85% of patients.
- On NRS scale, the mean varies between 3-6/10.
- Symptoms and severity in thalamic versus extrathalamic stroke does not differ.
- Intensity can be increased by internal or external stimuli.

Neurology 1995; 45: S11-S16.

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Spontaneous Pain Descriptions

- Continuous:
- Burning Aching Pricking
- Freezing Squeezing
- Intermittent:
- LaceratingShooting

- CPSP Can reduce quality of life:
- Can compromise rehabilitation.
 Interfere with sleep.
 Lead to self-mutilation.
- Even push patients to suicide.

ancet Neurol 2009; 8: 857-68

Pain Distribution

- Distribution of pain can range from a small area (eg, the hand) to large areas (eg, to one side of the body).
- •Large areas are most commonly affected, with or without involvement of the trunk and face.
- In patients with lateral medullary infarction, the pain can involve one side of the face and the contralateral side of the body or limbs, and periorbital pain is frequently reported.
- Hemibody pain is common in patients with thalamic lesions.

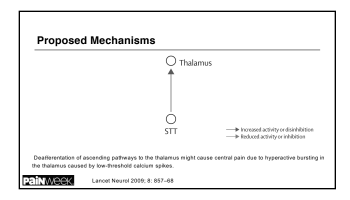


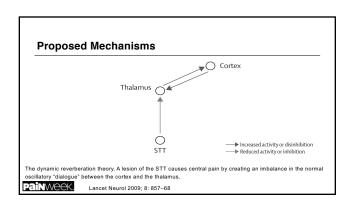
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Proposed Mechanisms Lateral Medial thalamus Medial thalamus Increased activity or disinhibition Reduced activity or inhibition Loss of STT input to the posterior lateral part of the thalamus causes disinhibition of the medial thalamus leading to pain. Painweek Lateral Medial thalamus Lateral Amerikanius causes disinhibition of the medial thalamus leading to pain.

Proposed Mechanisms Insula Posterior ventral medial nucleus Parabrachial nucleus periaqueductal grey STI The thermosensory disinhibition theory. A lesion in the lateral cool-signalling spinothalamocortical projections to the thermosensory area of the insula through the posterior part of the ventral medial incleus causes disinhibition of a medial limbic network involving the periaqueductal grey of the thermosensory area of the insula through the posterior part of the ventral medial incleus causes disinhibition of a medial limbic network involving the periaqueductal grey of the thermosensory area of the insula through the periaqueductal grey of the thermosensory area of the insula through the periaqueductal grey of the threaded. Painweck Lancet Neurol 2009; 8: 857–68

Proposed Mechanisms Lateral Medial thalamus Halamus Neospinothalamic/ Paleospinoreticulothalamic/ medial STT Increased activity or disinhibition A loss of normal inhibition from the rapidly conducting "neospinothalamic" or lateral STT projections causes disinhibition of the slowly conducting polysynaptic paleo spinoreticulothalamic or medial STT projections, resulting in pain. Painweek Lancet Neurol 2009: 8: 857–88





Antidepressants Anticonvulsants Antiarrhythmics Opioids	Regional Anesthesia Electrical Stimulation Deep Brain Stimulation Neuroablative Procedures
Steroids Intrathecal Baclofen Rehab Techniques	■ Neuroapiative Procedures ■ Transcranial Magnetic Stimulation
ainweek,	
Antidepressants	
■TCAs are currently viewed as	first-line drugs for CPSP.) is considered drug of choice, with consistent
	were common, particularly lethargy and dry
	oramine, desipramine) and otake inhibitors (venlafaxine, duloxetine, eported to be effective, but efficacies have yet to

Anticonvulsants

be established.

Pain Management Nursing 2015; 16(5): 804-818.
Pain 1989: 36: 27–36.
PEINWECK

Gabapentin and pregabalin have well documented efficacy in central neuropathic pain syndromes.

• Selective serotonin reuptake inhibitors are mostly ineffective.

- In a RCT, pregabalin showed a clinically significant effect of treatment on pain levels in patients with central neuropathic pain.
- Most commonly reported side-effects were dizziness, decreased intellectual performance, somnolence, and nausea.

Pain 2008; 136: 150–57.

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- Lamotrigine monotherapy was found to be moderately effective in amounts up to 200 mg/day in randomized double-blinded placebo-controlled trial of 27
- Lamotrigine was well tolerated except for the occurrence of mild rash. However, Stevens-Johnson syndrome and toxic epidermal necrolysis (TENS) are serious potential side effects of lamotrigine, and appropriate patient instruction must be given.

Am J Phys Med Rehabil. 2002;81(9):718-720.

Painweek.

Anticonvulsants

- In a placebo-controlled, crossover study comparing amitriptyline, carbamazepine, and placebo, carbamazepine was better at 3 weeks only, whereas amitriptyline was significantly better than placebo in relieving pain at 2, 3, and 4 weeks.
- ${\color{red} \bullet}$ Use of carbamazepine is limited by its side-effect profile and interaction with other medications.
- Clinicians should be aware of possible ataxia, rash, hyponatremia, bone marrow dysfunction, and hepatic dysfunction.

 Overall, the efficacy of carbamazepine is limited.

Pain 1989; 36: 27–36.

Opioids

- Opioids are generally considered ineffective in CPSP.
- However, morphine has been reported to alter significant aspects of pain perception (allodynia and thermal thresholds).
- In one study, morphine appeared to be effective in reducing CPSP because it reduced concurrent nociceptive pain and psychogenic influence.
- Other investigators have reported a loss or inactivation of opioid receptors in the cerebral hemisphere in CPSP, which would explain the low efficacies of opioids and the need for high doses to treat CPSP.
- Opioid treatment is often discontinued because of significant side effects from the high doses necessary for clinical benefit.

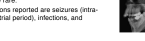
Pain Management Nursing 2015; 16(5): 804-818.

TABLE 3. Intravenous Drags Reported to be Effective in the Treatment of CPSP								
Drug	Reference	No. of Patients Total (CP)CPSP	Stady Design	Study	Desing Regimen	Outcome Measures	Resilix	Community
Lidocaine	Attail at af ²	16,06,6	Double-blind, placebe- coninsted, errosover	A	Singley IV over XV minutes	Spontanovas pain, VAS 1-100 Clobal assessment of pain solid	Sg greater relief of pain for up to 45 min with bloomine 11/15 > 59% pain relief with bloomine vs 6/15 with placebo	Also sig reduction of bendu- induced allindysis and mechanical hyperalgesia
Propoful	Canaziero et al ^{ne}	32,16/7	Double-blind, placebe- controlled, erresoner	A	Single IV bolas of 0.2mg/kg respondens 0.3mg/kg per k for 6-24 h	VAS 1-10 every 5-min for 30-min	Reduction by >3 VAS points in 5/7 CPSP patients, 14/16 CP parients vs 6/16 with placebo	In responders all of paid abeliabed, pain control with prolonged inflation for 6-24 h in 7 parients
Keiamine	Buckenja ci al ^e	632	Double-blind, placebo- controlled, errne-over	Α	256yg kg IV ever 5 min	Pain rating scale 0-10	Pain relief > 90% in 2/3 patients with CP (both with CPSP) busing 2-3 beam to 9/3 with placeby	Continues subcutaneous influsion only in I patient with neuropathic pain, discontinued because of side-offices
	Vanameto et al	23(23(23	Uncontrolled trial dl morphise d2 thiampful d9 lecumine	я	Sing every Sinin, total dose 25mg	VAS 1-10	Pain relief > 40% in 11/23 patients, 2/23 pain increase, datation < 98 min	Ne long-term application
Thiospial	Yamameto et al ²⁰	28/29/29	Uncontrolled trial, dl morphine, dl thiamplal, dl latamine	В	50 mg every 5 min, total desc 250 mg	VAS 1-30	Pain relief > 40% in 22/29 patients, duration > 60 min	No long-term application
Morphine	Yamamoto et al ²⁰	29,39,29	Uncontrolled trial, dl morphine, d2 thiomplel, d1 letterine	в	Jung every 5 min, total dose 18 mg	VAS 1-10	Pain odid" > 40% in 839 patients, duration > 60 min	No long-term application
	Anal et al ⁷⁹	15/15/6	Planto-controlled, conserver	A	Mean design Hing IV Mean design 97mg ceil	VAS 1-100	Ne sig difference in pain reduction 4 of 14 with long-term officacy of onal morethine	Sig influence of morphise on allodysis and thermal threshold

Neurostimulation

- Motor cortex stimulation:
- —Mechanism not completely understood. However, studies have indicated changes in cerebral blood flow in several areas, including the thalamus, after successful motor cortex stimulation.

- -In two recent reviews, the 1-year success rate in patients with CPSP was concluded to be about 45-50%.
- -Severe complications are rare.
- Most common complications reported are seizures (intra-operatively or during the trial period), infections, and hardware problems





Lancet Neurol 2009; 8: 857–68

Neurostimulation

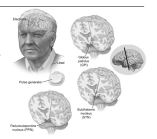


- Transcranial magnetic stimulation:
- -Non-invasive method.
- -The effects on pain are often modest and short lasting.
- -Adverse events are rare.
- Recurring sessions of repetitive transcranial magnetic stimulation of the motor cortex have been shown to extend pain relief.
- -The result of this treatment might be a useful predictor for the efficacy of motor cortex stimulation.

Lancet Neurol 2009; 8: 857–68

Neurostimulation

- Deep brain stimulation:
 - Main targets are the sensory (ventral posterior) thalamus and the periventricular gray matter.
 - -Reported efficacy rates range from 25% to 67%, but with wide ranges of pain relief.



Lancet Neurol 2009; 8: 857-68

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Neurostimulation



- Vestibular caloric stimulation:
 - -Effect probably due to activation of the posterior insula and subsequent inhibition of pain generation in the anterior cingulate.

 -Two small studies:

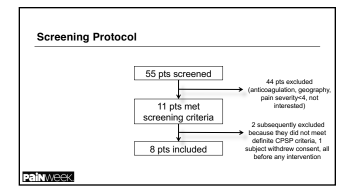
 - -In one study (n=2), CPSP was substantially relieved by VCS.
 - In another study of 9 patients, there was a significant immediate treatment effect for cold-water caloric stimulation.

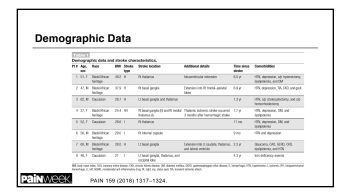
Neurocase 2007; 13(3): 185-188.

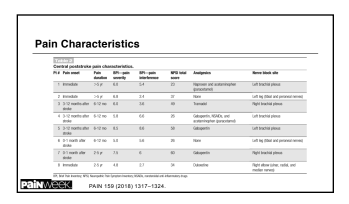
Journal of Neurology Neurosurgery and Psychiatry 2008; 79(11): 1298-1301.

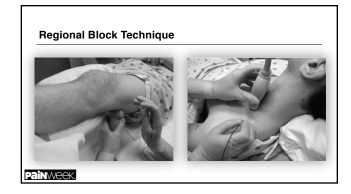
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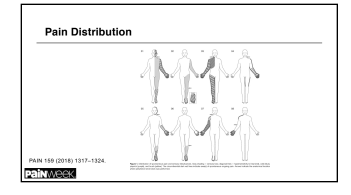
How central is central poststroke pain? The role of afferent input in poststroke neuropathic pain: a prospective, open-label pilot study

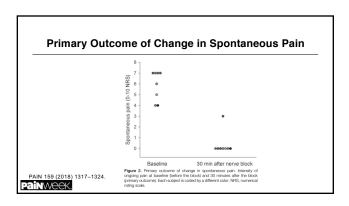


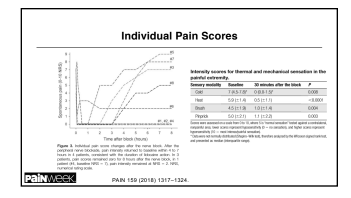


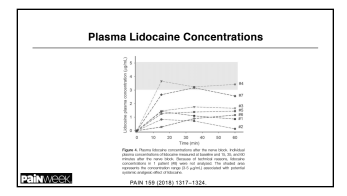












Discussion

- Pain may not be entirely generated and perceived in the CNS.
- Rather, the afferent sensory input from the painful area plays a role in maintaining spontaneous pain in CPSP.
- It is plausible that the sensory neurons in the CNS, which are damaged by the stroke, become sensitized to the afferent stimuli, and generate action potentials secondary to trivial sensory input.
- Supporting the local afferent blockade (rather than the systemic effect) as the cause of pain relief is the finding that no changes in pain intensity occurred after the block in the ipsilateral painful extremity in these patients.

PAIN 159 (2018) 1317-1324.

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- CPSP has a variable time to onset after stroke.
- In most cases of CPSP, the stroke lesions are extrathalamic.
- Amitriptyline is the first-line drug of choice.
- If amitriptyline fails or is unavailable, then try lamotrigine.
- In intractable cases, short-term pain relief may be achieved by IV lidocaine, propofol, or ketamine.
- Motor cortex stimulation, DBS, or, rTMS may be tried in resistant CPSP patients.
- Sensory afferent input may play an important role in maintaining pain in CPSP.

PAIN 159 (2018) 1317–1324.