Introduction

The 2019 Annual Meeting of the Eastern Pain Association provided a clear foundation for the future of pain medicine as it heralded the year 2020. The best way to optimally manage chronic pain patients of the future is for pain professionals to go back to our past as a multidisciplinary discipline. The multimechanistic, multimodal and multidisciplinary approach lies at the heart of Eastern Pain Association’s mission as a multidisciplinary society that is dedicated to disseminating scientific knowledge as it relates to pain medicine through improved pathophysiological understanding, discovery of new therapeutic targets and interventions to improve patient outcomes.

Moderation and a combination approach are key. We are not allowing the field of pain to be reduced to pro vs anti-opioid or pro vs anti-interventional pain wars. The focus is to gain a deeper understanding of our patients’ pain through improved knowledge, and the abstracts presented at the Eastern Pain Association’s 2019 Annual Meeting are a small step in that direction.

The three award-winning abstracts that are presented here give us a glimpse into what the future holds, from improved understanding of chronic pain mechanisms to further progress in the field of regenerative medicine. Enjoy.

1. Spinal Cord Microglial Phenotypic Changes Following Sciatic Nerve Crush in CD137LKO Mice

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Purpose: Neuropathic pain is defined as pain caused by a lesion or disease of the somatosensory system. This debilitating illness affects roughly 7 percent of the population in the United States. Microglia, important immune cells in the central nervous system, are shown to play a critical role in the development of neuropathic pain. Peripheral nerve injury can activate spinal cord microglia to become pro-inflammatory versus anti-inflammatory phenotypes. CD137 ligand is a receptor on microglia that binds to the CD137 receptor on T lymphocytes. We have shown that CD137L knockout (KO) mice display reduced sensory sensitivity and faster functional recovery following sciatic nerve crush (SNC). We hypothesize that CD137L depletion induces this change through promoting microglia to preferentially differentiate into an anti-inflammatory subtype after SNC.

Methods: To test this hypothesis, qRT-PCR and Flow Cytometry were used to measure the expression of markers specific to pro- vs anti-inflammatory microglia at various times following SNC in both wild-type (WT) and CD137L KO mice.
**Results:** Data from qRT-PCR showed that CD137L KO microglia displayed a more pronounced increase of IRF3 and earlier induction of downstream chemokines CCL3, CCL4 and CCL5 compared to WT microglia. The CD137L KO microglia also showed delayed or lack of upregulation of CD86 and MyD88-downstream chemokines, CCL2 and CXCL10 following SNC compared to WT microglia. Preliminary flow cytometry data with WT mice showed increased total microglial and CD86+ microglial numbers within lumbar spinal cord post-SNC.

**Conclusion:** Our data so far appear to support the prediction that microglia from CD137L KO mice exhibit an increased anti-inflammatory profile compared to WT mice following SNC injury. CD137L and its downstream pathways could serve as potential drug targets.

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**2. Notalgia Paresthetica Successfully Treated with Cervical Epidural Injection and Occipital Nerve Block: A Case Report**

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**Background:** Notalgia paresthetica (NP) is a common cutaneous dysesthesia involving the dorsal spinal nerves that is largely under-recognized and underdiagnosed. It is characterized by a unilateral (usually) neuropathic pruritus normally occurring in the interscapular and paravertebral region resulting in pain, discomfort, scarring from incessant itching and decreased quality of life. NP is usually localized to the T2–T6 dermatomal distribution and accompanied by localized burning, tenderness, hyperalgesia and dysesthesia resulting in chronic itching and a post-inflammatory hyperpigmented patch that is esthetically distressing.1 The etiology of NP is not fully understood; however, it is widely accepted that the sensory neuropathy is caused by musculoskeletal compression of the cutaneous branches of the dorsal primary rami of thoracic spinal nerves T2–T6.1–2 While NP is associated with symptoms in the T2–T6 dermatomes, it has been associated with cervical spinal disease usually at the C4–C6 level even though the spinal pathology seen on imaging does not correlate with the dermatomal distribution of symptoms.2–6 Current first-line therapies include topical treatments such as lidocaine and capsaicin, conservative management with physical therapy, manual manipulation and/or transcutaneous electrical nerve stimulations (TENS) and medications such as gabapentin with varying and/or temporary success. Surgical and minimally invasive interventions are reserved for refractory cases and are rarely pursued. In this article, we present a case of NP that was successfully treated with a cervical interlaminar epidural steroid injection (ESI) and bilateral occipital nerve block, modalities typically used for cervical spinal pathology and cervical neuralgias, respectively, but not previously described for NP.

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Notalgia paresthetica (NP) is a common cutaneous dysesthesia involving the dorsal spinal nerves that is largely under-recognized and underdiagnosed. It is characterized by a unilateral (usually) neuropathic pruritus normally occurring in the interscapular and paravertebral region resulting in pain, discomfort, scarring from incessant itching and decreased quality of life. NP is usually localized to the T2–T6 dermatomal distribution and accompanied by localized burning, tenderness, hyperalgesia and dysesthesia resulting in chronic itching and a post-inflammatory hyperpigmented patch that is esthetically distressing.1 The etiology of NP is not fully understood; however, it is widely accepted that the sensory neuropathy is caused by musculoskeletal compression of the cutaneous branches of the dorsal primary rami of thoracic spinal nerves T2–T6.1–2 While NP is associated with symptoms in the T2–T6 dermatomes, it has been associated with cervical spinal disease usually at the C4–C6 level even though the spinal pathology seen on imaging does not correlate with the dermatomal distribution of symptoms.2–6 Current first-line therapies include topical treatments such as lidocaine and capsaicin, conservative management with physical therapy, manual manipulation and/or transcutaneous electrical nerve stimulations (TENS) and medications such as gabapentin with varying and/or temporary success. Surgical and minimally invasive interventions are reserved for refractory cases and are rarely pursued. In this article, we present a case of NP that was successfully treated with a cervical interlaminar epidural steroid injection (ESI) and bilateral occipital nerve block, modalities typically used for cervical spinal pathology and cervical neuralgias, respectively, but not previously described for NP.

**Case Report**

A 46-year-old female presented to clinic with severe pruritus in her bilateral periscapular and occipital region for 1 year (Figure 1). The pruritus started in the neck and progressed cephalad and caudad. She had accompanying neck pain with...
occasional radiation down the left arm at the time of symptom onset and was presumed to have cervical radiculopathy. Her pain mostly resolved with a Medrol dose pack and a course of physical therapy, but pruritus persisted. On initial visit, physical exam demonstrated mild pain with cervical rotation at end range of motion, negative Spurling’s, but with scattered scarring across the medial scapular region bilaterally. She was prescribed topical anesthetic gel/ointment and gabapentin with no relief; her pruritus remained persistent and constant. The patient was referred to dermatology where she was diagnosed with suspected allergic reaction with residual irritation and prescribed 4 weeks of Allegra which conferred minimal relief. Imaging was obtained with cervical X-ray showing evidence of disc space narrowing at C5–6. Follow-up cervical MRI showed disc bulging at C4–5, C5–6. The patient subsequently underwent an interlaminar C7-T1 ESI using 1 mL of Depo-Medrol with near-complete resolution of pruritus in the scapular region but ongoing in the occipital region (Figure 2). She later underwent two occipital nerve blocks using 3 mL of 1% lidocaine and 0.25 mL celestone per side with 70% improvement in symptoms.

**Discussion**

There are currently numerous treatment alternatives for NP; however, the majority only confer temporary symptomatic relief either losing efficacy over time or with cessation of treatment. Topical treatments include capsaicin, tacrolimus and anesthetic creams. Intralesional treatments include botulinum toxin A, corticosteroids (triamcinolone) and cryolipolysis. Oral medications include oxcarbazepine, gabapentin and amitriptyline. Other therapies documented include TENS, EMS, Narrow band UV-B, osteopathic manipulation, acupuncture, exercise and physiotherapy. Paravertebral nerve blocks,
intravenous lidocaine infusions\textsuperscript{27} and surgical decompression\textsuperscript{28} have also been described. ESI and occipital nerve block have not yet been described as viable treatments for NP. In order to understand why these interventions succeeded in conferring relief, we must acknowledge the following precepts: 1) pruritus can be a symptom of nerve damage with neuropathology as opposed to dermatopathology and 2) NP results from damage to the cutaneous branches of the dorsal primary rami of thoracic spinal nerves T2–T6 by compression or impingement from degenerative changes in the spine or entrapment from paraspinal muscle spasms.\textsuperscript{29} In 1979, Massey and Pleet suggested that, specifically, the posterior rami of spinal nerves T2–T6 are more susceptible to nerve entrapment and chronic trauma because they pass through the multifidus spinae muscle at a 90-degree angle.\textsuperscript{21,30} This theory of muscle spasms/entrapments coincides with the success of treatments such as physiotherapy which takes into account the anatomy of paraspinal muscles in relation to the spinal nerves. The upper six thoracic spinal nerves first pierce the rhomboid and trapezius prior to becoming cutaneous nerves. Fleischer et al used this knowledge to develop an exercise which involved strengthening the rhomboids and latissimus dorsi muscles and stretching the pectoralis muscles to reduce the angle of the nerve as it passes through the muscles. Extension of the spine through particular exercises may also reduce the angle of the nerve at the spinal level and thus the pressure it experiences.\textsuperscript{25}

The second school of thought posits that nerve impingement and compression of the upper thoracic spinal nerves is caused by cervicothoracic musculoskeletal degenerative processes; the nerves are entrapped as they exit the spine through the vertebral foramen. One could argue that the characteristic pruritus experienced is located along the T2–T6 dermatomes and does not correspond with a pathological cervical dermatomal distribution. Facet arthropathy, however, can present as referred pain in the upper thoracic and infrascapular region, with symptoms from the cervical area referred directly to the infrascapular back which is the characteristic site of pruritus in NP (Figure 3). While degenerative vertebral disk disease corresponding to the affected dermatome may also be observed, degenerative joint disease in the cervical region resulting in pain in a thoracic dermatome should not be discounted; through clinical observations, there exists a clear relationship between the upper thoracic region and the cervical spine.\textsuperscript{31}

In 2000, Savk and Savk demonstrated that of 10 patients with NP, 7 showed evidence of degenerative changes in the vertebrae. While most changes were most prominent in the vertebrae that corresponded with the dermatomal cutaneous lesion, the study did demonstrate cervicothoracic disease corresponding with T4–T8 NP localization.\textsuperscript{5} In 2005, they evaluated 43 patients with NP and found that 79.1\% of the cohort had positive radiographic findings consistent with degenerative changes or herniated nucleus pulposus in the cervicothoracic region.\textsuperscript{32} Other studies have also shown a correlation between spinal pathology and NP.\textsuperscript{3,20} As described earlier, the patient continued to have occipital and upper cervical pain even after the cervical ESI. This is likely because the upper cervical spinal pathology noted on imaging was not adequately penetrated with the C7-T1 ESI. In addition, repetitive upper body and neck twisting associated with incessantly scratching ones back likely contributed to a concomitant cervicalgia. It is likely that the occipital nerve block conferred near-complete resolution of neck pain and irritation by accessing the upper cervical

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{Figure3.png}
\caption{Facet Mediated Referred Pain.}
\textbf{Notes:} Depiction of referred pain of the cervical neck region. Note that the lower cervical region pain referral pattern overlaps onto the scapular region and the upper thoracic region.
\end{figure}
region as well as providing anesthetic relief to cervical muscle strain. It is also possible that the neck and occipital pain experienced was a superimposed atypical occipital neuralgia.

Conclusion

Notalgia Paresthetica, while defined as sensory neuropathy, may also be a cutaneous sign of underlying degenerative cervical or cervicothoracic disk disease. This is supported, in this case, by the significant improvement of scapular pruritis, in a patient with underlying cervical disk disease, who received a cervical ESI. In patients who present with NP, cervical spinal imaging is appropriate, and first-line treatment should include addressing the underlying spinal pathology. Topical and/or conservative measures will likely only confer temporary relief.

3. Evaluation of Umbilical Cord derived Wharton’s Jelly for Regenerative Medicine Applications

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Introduction: Musculoskeletal injuries have traditionally been treated with activity-modification, physical therapy, pharmacological agents and surgical procedures. These modalities have limitations and potential side-effects. Over the last decade, there has been an increased interest in the use of biologics for regenerative medicine applications (RMA), including umbilical cord (UC) derived Wharton’s Jelly (WJ). Despite this increase, there is insufficient literature assessing the amount of growth factors, cytokines, hyaluronic acid (HA) and extracellular vesicles (EV) including exosomes in these products. The purpose of this study was to develop a novel WJ formulation and evaluate the presence of growth factors, cytokines, HA and EV including exosomes.

Materials and Methods: Human-UC was obtained from consenting C-section donors. The WJ was then isolated from the procured UC and formulated into an injectable form. Randomly selected samples from different batches were analyzed for sterility testing and presence of growth factors, cytokines, HA, and particles in the EV size range.

Results: The results revealed that all samples passed the sterility test. Growth factors including IGFBP 1, 2, 3, 4 and 6, TGF-α, PDGF-AA were detected. Expression of several immunomodulatory cytokines, such as RANTES, IL-6R, IL-16 were also detected. Expression of pro-inflammatory cytokines MCSFR, MIP-1a; anti-inflammatory cytokines TNF-RI, TNF-RII, IL-1RA; and homeostatic cytokines TIMP-1 and TIMP-2 were observed. Cytokines associated with wound-healing, ICAM-1, GDF-15, and regenerative properties, growth hormone, were also expressed. High concentrations of HA were observed. Particles in the EV size range (30–150 nm) were detected and were enclosed by the membrane, indicative of true EV.

Conclusion: Our results confirmed there are numerous growth factors, cytokines, HA and EV present in the WJ formulation we analyzed. We believe the presence of multiple factors within one WJ formulation may play a role in reducing inflammation, pain and augment healing of musculoskeletal injuries. This offers a potential expanded use for RMA.

Keywords: regenerative medicine, umbilical cord, Wharton’s jelly, growth factors, cytokines, exosomes

References