However, disease activity did not differ significantly in either group (group I 5.6 ± 0.6 vs 4.8 ± 1.6 p = 0.4; group II 4.9 ± 0.3 vs 5.2 ± 1.2 p = 0.8).

Conclusion: This preliminary data shows greater effectiveness of Pregabalin in comparison with NSAID and DMARD treatment, both in terms of pain intensity and the neuropathic pain component, which is of practical importance.


THU0487 PERINEURAL INJECTION THERAPY; A NEW MODALITY IN MANAGEMENT OF MECHANICAL LOW BACK PAIN; A COMPARATIVE STUDY

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Background: Chronic mechanical low back pain represents the second leading cause of disability worldwide being a major welfare and economic problem. Subcutaneous prolotherapy treats prolonged pathological peripheral neurogenic inflammation for several painful conditions. As it induces apoptosis of proliferating peptidergic nociceptors and neovessels by reducing vascu lar endothelial growth factor. Objectives: To assess the effectiveness of subcutaneous perineural injection therapy in management of pain, physical function, disability and psychological status in mechanical low back pain

Methods: Ninety patients with non-radiating non-specific chronic mechanical low back pain (LBP) that persisted for 12 weeks or more were selected in this study. (Patients with inflammatory LBP, radiating, or LBP due to specific cause, pregnant women or Patients with implanted pacemaker or spinal cord stimulator were excluded from this study). After giving written consent; the patients were randomly divided into 3 groups 30 patients in each group. Group I received 8 weekly subcutaneous injections of 1 ml of buffered dextrose 5% in each chronic constriction injury points and tender points in back and buttock, Group II treated by using pulsed electromagnetic field stimulation (PEMFs) for 30 minutes over the lower lumbar region with frequency of 10 Hz and intensity of 2 mT/milltesla every other day for 8 weeks, Group III received sham PEMFs. All patients will be instructed to follow an exercise program in the form of stretching for other day for 8 weeks. Group III received sham PEMFs. All patients will be instructed to follow an exercise program in the form of stretching for another day for 8 weeks. Group III received sham PEMFs. All patients will be instructed to follow an exercise program in the form of stretching for another day for 8 weeks. Group III received sham PEMFs. All patients will be instructed to follow an exercise program in the form of stretching for another day for 8 weeks. Group III received sham PEMFs. All patients will be instructed to follow an exercise program in the form of stretching for another day for 8 weeks.

Results: Mean VAS was 7.9 ± 1.4, 8.2 ± 0.6, and 7.6 ± 1.1 respectively. No baseline differences existed between all groups in all parameters. There was significant improvement in VAS, WOMAC and HDAS in group I & II (p<0.05) after treatment, and 4 months later (figure 1). While the improvement in group III was non-significant (p>0.05). The improvement in group I was better than in group II with significant difference between the two groups (p<0.05) after treatment and 4 months follow up.

Conclusion: Perineural Injection Therapy is an effective new modality in management of pain, physical function, psychological status and disability in mechanical low back pain.

REFERENCES:


THU0488 ACHILLES TENDON RUPTURE ASSOCIATED WITH THE USE OF FLUOROQUINOLONES IN PATIENTS OLDER THAN 60 YEARS

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Background: Fluoroquinolones (FQ) are a class of broad-spectrum antibiotics whose use has spread as they are considered safe and well tolerated drugs. Among its musculoskeletal side effects, Achilles tendon rupture (ATR) is a well-known complication that can be disabling, arising recent interest from the pharmacovigilance system after evaluating the reported side effects.

Objectives: To describe the epidemiological and clinical features of patients diagnosed with FQ-associated ATR in a Spanish tertiary hospital.

Methods: A retrospective observational study was performed, which included all patients older than 60 years who were diagnosed of ATR in our center during the period 2000-2017, identifying patients who had been previously treated with FQ. The demographic, clinical and outcome data were obtained from their medical records.

Results: During the study period, 44 patients with ATR were identified, 8 (14.6%) of them previously treated with FQ. In this group of patients, the mean age at diagnosis of ATR was 77.37 ± 9.54 years, being male 6 (75%). Four of them (50%) received concomitant treatment with corticosteroids (CS) and one patient had undergone kidney transplantation due to nephroangiosclerosis. Seven patients (87.5%) were treated with Levofloxacin and one case received Ciprofloxacin, all of them orally. The indication for FQ treatment in half of the cases was acute bronchitis and in the other half exacerbations of underlying respiratory pathology (chronic obstructive pulmonary disease and diffuse interstitial lung disease). The mean duration of treatment with FQ was 6.16 ± 2.4 days, while the mean time from the start of treatment to the diagnosis of ATR was 19.95 ± 14.83 days. In seven patients (87.5%) the rupture was spontaneous, while one patient presented traumatic rupture (low impact traumaism), 87.5% of the ruptures were total ruptures and all cases required surgical treatment, without recurrence reported. The comparison of the characteristics of patients with ATR who had or not received treatment with FQ is shown in the table, identifying significant differences in favor of a higher percentage of patients who were smokers, received concomitant treatment with CS and had spontaneous rupture in the group of patients who had received FQ.
Conclusion: The Achilles tendon is the most frequent location of tendinopathy associated with FQ, being affected in 95% of cases. The risk factors associated to an increased risk to develop FQ-associated ATR includes age over 60 years, male gender, chronic treatment with CS and organ transplantation, all these being present in our cases. Despite being a relatively frequent adverse event, it is underdiagnosed and the risk of ATR is not usually assessed when indicating FQ treatment. It is important to perform a risk/benefit assessment, specially in patients with associated risk factors, because most ruptures are complete and require surgical treatment, and may be a potential cause of disability.

REFERENCES:

Disclosure of Interests: None declared

THU0490 PREDICTORS OF CHRONIC PAIN RELIEF BY FIXED-SITE HIGH-FREQUENCY TRANSCUTANEOUS ELECTRICAL NERVE STIMULATION
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Background: Fixed-site high-frequency transcutaneous electrical nerve stimulation (FS-TENS) is a form of TENS in which the stimulator is designed for a predetermined location rather than for co-localization with the patient’s pain. Previous studies in individuals with chronic pain have demonstrated the efficacy and real-world effectiveness of a small wearable FS-TENS device designed for localized application to a single placement site to ensure comfort while both active and sleeping. Results demonstrated that 50–80% of FS-TENS users with chronic lower extremity or low back pain experience clinically meaningful pain relief, and daily use of the device was associated with greater pain relief relative to intermittent use.

Objectives: To determine predictors of a positive FS-TENS response.

Methods: This retrospective, observational study evaluated users of a FS-TENS device to treat chronic pain over a 10-week period. The device and companion smartphone app collected dosage data, demographics, pain characteristics, and pain ratings and all data were stored in a cloud database. The primary study outcome was the baseline to 10-week change in composite pain (average of pain intensity and pain interference with sleep, activity, and mood). Device users were included if they provided demographic data, pain characteristics indicative of chronic pain, and baseline and week 10 pain ratings. Participants were defined as a responder or comparator based on their change in composite pain (responder: >15% decrease; comparator: >15% increase). Stepwise forward probit regression was used to determine independent predictors.

Results: There were 451 responders and 263 comparators. Independent predictors (Table) that were associated with greater response to FS-TENS included age, baseline composite pain, adherence/utilization rate (defined as the percentage of days with at least 30 minutes of stimulation), and stimulation intensity (defined as the ratio of stimulation to sensation threshold, expressed in decibels). Negative predictors (associated with lower response) included history of headache/migraine and diabetes. The area under the receiver operating characteristic curve was 0.76 (95% confidence interval, 0.72–0.79).

Conclusion: FS-TENS effectiveness is predicted from baseline pain characteristics and dosage variables, such as frequency/regularity of use and stimulation intensity, with moderate accuracy.

REFERENCE:

Abstract THU0490 – Table 1. Independent predictors of responders from probit regression.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coefficient</th>
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<tbody>
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<td>Age (years)</td>
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<tr>
<td>Baseline composite pain (0–10)</td>
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<tr>
<td>Headache or migraine</td>
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