Comparison of venlafaxine and duloxetine: measuring clinical impact of time to therapeutic dose (TTD) among patients achieving therapeutic dosing for pain

Abstract

Background: Chronic opioid therapy remains controversial; however, there is consensus among treatment guidelines that adjunct medications should be utilized first. A meta-analysis revealed serotonin-norepinephrine reuptake inhibitors (SNRIs) venlafaxine and duloxetine to be equally efficacious in the treatment of neuropathic pain and recommend them as first-line therapy. Although, the therapeutic dosing for pain with both medications is well established there are no head-to-head studies comparing the two medications. The impact of medication selection (TIMS) as measured by percentage of patients achieving therapeutic dose and time to therapeutic dose (TTD) for neuropathic pain with either venlafaxine or duloxetine is not well understood. These outcomes along with adverse effect profiles need to be evaluated to inform clinical decision making.

Materials and Methods: This was a single center, retrospective, observational analysis. New start prescriptions for either venlafaxine or duloxetine between January 1, 2011 and January 1, 2014 were identified. Through data warehouse extraction the following was collected: age, gender, weight, height, race, comorbidities, prescriber and concomitant antidepressants and anticonvulsants on date of initiation. Manual data collection through the Computerized Patient Record System (CPRS) was then utilized to determine veteran eligibility as well as if therapeutic dose was achieved, TTD, as well as discontinuation rates and cause.

Results: 682 charts were reviewed to identify 302 patients, 151 in each group. The duloxetine group had 120 patients achieve therapeutic dose compared to 82 in the venlafaxine group (p<0.0001). Median TTD for duloxetine was 7 days (0-44.25, IQR) compared to venlafaxine 31.5 days (10-115, IQR). At study conclusion, 50/151 (33.1%) patients remained on duloxetine compared to 31/151 (20.5%) of those on venlafaxine (p-value 0.0191). Side effects were reported in 37% of patients in venlafaxine group compared to 22% of duloxetine group (p=0.0053). Of note, 117 (77%) of the duloxetine patients had a previous trial of venlafaxine therapy.

Conclusion: Patients taking duloxetine are significantly more likely to achieve therapeutic dose, arrive at therapeutic dose more quickly, and remain on the medication compared to venlafaxine. Titration schedule may influence tolerability. Duloxetine should be favored over venlafaxine in treatment algorithms for neuropathic pain.

Keywords: antidepressants, duloxetine, venlafaxine, neuropathic pain, diabetic neuropathy, pain

Introduction

Managed Care and integrated health systems have obligations to control costs in their allocation of limited resources and utilize formulary management to balance cost savings concerns with available evidence. Pharmacoeconomic evaluations often decide the availability of therapies based purely on cost when no evidence exists for clinical superiority between two treatments. From a therapeutic perspective, this presents challenges when multiple medications are recommended as first-line options in available guidelines but no head-to-head studies exist to establish superiority. The cost of randomized controlled trials (RCTs) to establish such superiority is prohibitive and therefore they are rarely performed. Randomized trials are also not typically reflective of clinical practice as they exclude higher risk or medically complex patients and do not last long enough to evaluate long-term efficacy, rate of discontinuations, and number of patients that remain on subtherapeutic doses. Patient’s, however, do have the right to the most effective treatment and delaying or diverting treatment away from effective therapy for cost considerations has significant ethical considerations. Neuropathic pain is often a chronic, debilitating condition, with a complex pathophysiology. The American Academy of Pain Medicine highlighted a recent market research report which indicates that more than 1.5 billion people worldwide suffer from chronic pain and that approximately 3-4.5% of the global population suffers from neuropathic pain, with incidence rate increasing in the aging population. Current evidence-based treatment guidelines recommend the use of amitriptyline, duloxetine, venlafaxine, gabapentin, or pregabalin as first line therapy.
Despite the efficacy of tricyclic antidepressants (TCAs) being largely established, they have moved out of favor due to their excessive side effects which include dry mouth, sedation, and blurred vision.1,2 Serotonin norepinephrine reuptake inhibitors (SNRIs) have largely replaced TCAs, working through the same pathway by inhibiting reuptake of norepinephrine, and are better tolerated.3 Venlafaxine and duloxetine, both SNRIs, have been shown to be effective in RCTs.3,9-11 Clinical trials have shown that neuropathic pain relief by SNRIs is dose dependent which is heavily influenced by receptor selectivity. Duloxetine is highly selective for norepinephrine with a receptor affinity (10:1) compared to venlafaxine (30:1), respectively.12 Therefore, venlafaxine requires a higher dose to reach analgesic therapeutic effect unlike duloxetine. This was reflected in clinical trials when efficacy for venlafaxine was shown at doses greater than 150mg/day and duloxetine 60mg daily.13,14 Literature assessing time to therapeutic effect of venlafaxine and duloxetine in the treatment of pain is limited to pharmaceutical trials; no head-to-head studies have been conducted. As venlafaxine and duloxetine have very different titration schedules, the difference in tolerability between medications depending on how quickly each is titrated to their therapeutic dose may play a large role in therapeutic success. In addition to percentage of patients achieving therapeutic dosing and time to therapeutic dose, the percentage of patients that remained on the medication would be very useful as long-term efficacy is not typically assessed in clinical trials. To address the above concerns we elected to perform a therapeutic impact of medication selection (TIMS) evaluation between venlafaxine and duloxetine which are both first-line options recommended for neuropathic pain.

**Methods**

This is a single center, retrospective, observational analysis that was conducted at the Tennessee Valley Healthcare System (TVHS), an integrated Veterans Affairs (VA) system consisting of two medical centers and several community-based outpatient clinics located throughout middle Tennessee and southern Kentucky. The study protocol was approved by TVHS Institutional Review Board (IRB) and exempted from patient notification because it was retrospective, observational, and de-identified. Using the Veterans Integrated Service Network 9 (VISN 9) data warehouse, an initial patient list was pulled identifying patients who were receiving care at TVHS between January 1, 2011 and January 1, 2014 with a new start prescription for either venlafaxine or duloxetine. Patient records that included previous trials of the same SNRI within a 5 year period and/or were being prescribed venlafaxine or duloxetine for an unspecified or non-pain related indication were excluded. Additional data pulled included comorbidities, concomitant antidepressants by pharmacological class on date of initiation, active prescription for gabapentin or pregabalin on date of initiation, most recent eGFR at date of initiation, and patient demographics.

As to time to therapeutic dose has not been assessed in a head to head study for these medications, we used discontinuation rates as a surrogate marker to determine minimum sample size needed. Based off of clinical trials this study required 302 patients, 151 patients per arm, to have at least 80% power allowing for detection of a difference of 15 percent increase or decrease of patient’s titrated to therapeutic dose between venlafaxine and duloxetine treatment groups. After the patient list was compiled, manual data collection was utilized to determine if patients were eligible for inclusion. Patients were included in the analysis if they were ≥18 years of age, eGFR >30, prescribed a new start prescription for venlafaxine or duloxetine with documented indication for pain within the specified time period. Patients were excluded if they had a previous trial of the same SNRI within a 5-year period and/or are being prescribed venlafaxine or duloxetine for unspecified or non-pain related indication. Those patients identified as fitting the inclusion criteria were then subject to a more extensive chart review. Starting dose and current dose were collected, as well as the difference in number of days from initiation to therapeutic dose or discontinuation. If SNRI prescription was discontinued or expired, chart notes were reviewed to determine cause.

All analyses were performed using the Excel or GraphPad Software systems. The Fisher’s Exact was test was used to compare rates of discontinuation, percent of veterans reaching therapeutic dose and all baseline demographics, except age. Age was assessed using the t-test. Nonparametric or non-continuous data is described using median and interquartile ranges and statistical analysis using Mann-Whitney U. Statistical significance was set at p<0.05 and all tests were 2-tailed (Figure 1).

![Figure 1](image)

**Figure 1** Percentage of patients achieving therapeutic dose.

**Results**

A total of 682 charts were reviewed with 302 patients meeting inclusion criteria, and 151 patients in each of the venlafaxine and duloxetine groups respectively. Baseline demographics were equivalent between groups, except for age, which was higher in the duloxetine group (55) versus venlafaxine group (48) and appears statistically significant (p-value<0.0001). For the primary endpoint of percentage of patients achieving therapeutic doses (TD), the duloxetine group had 120/151 (79.5%) versus the venlafaxine group which had 82/151 (54.3%) (p-value<0.0001). For the primary endpoint of time to TD, the median time for duloxetine was 7 days (IQR 44 days) versus median time for venlafaxine was 31.5 days (IQR 105 days) (p-value 0.0001) (Figure 2).

For the secondary outcomes, the percentage discontinuation rate for duloxetine was 101/151 (66.9%) with 50/120 (41.7%) remaining on duloxetine at study conclusion compared to 120/151 (79.5%) with 31/82 (37.8%) remaining on venlafaxine. In absolute numbers, 50/151 (33.1%) patients started on duloxetine remained on it at study conclusion compared to 31/151 (20.5%) of those started on venlafaxine (p-value 0.0191). The percentage discontinuation rate (Table 1) based on speed of titration for duloxetine versus venlafaxine showed, for <14 days to TD 64.5% compared to 87.5%, for 14-30

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days to TD 55.6% compared to 80%, for 30-60 days to TD 55.6% compared to 90%, for 60-90 days to TD 25% compared to 50%, and for >90 days to TD 63.6% compared to 48.3% (Figure 3). Reasons for discontinuation prior to achieving therapeutic dosing were primarily change in mood (13.25% vs 9.27%), GI distress (4.64% vs 4.64%), sexual dysfunction (3.97% vs 2.65%), sedation (2.65% vs 2.65%), and nightmares (2.65% vs 1.32%) for venlafaxine and duloxetine respectively (Table 2). Concomitant diseases and medications between groups is represented in Table 3. Out of the 151 patients with new start prescriptions for duloxetine, 117 (77.48%) tried venlafaxine prior to duloxetine. Of the 117 patients who trialed venlafaxine first, only 70 (59.83%) achieved therapeutic dose. 35 of the 117 patients (29.91%) received one prescription for venlafaxine before discontinuation (Table 1)(Figure 3).

**Discussion**

Chronic opioid therapy remains controversial and its appropriateness, in chronic pain management, a subject of national debate. There is consensus, however, in treatment guidelines that adjunct medications should be utilized first and there is a sense of urgency to provide effective relief for patients in a timely manner. Lack of efficacy and intolerable side effects complicate treatment with adjunct medications and represent a significant source of frustration for patients. The therapeutic impact of medication selection (TIMS) is a critical concept to clinical practice and specifically pain management to determine which first-line therapy has the highest chance of efficacy, shortest time to therapeutic dose, and best long-term success. We applied this evaluation to use of venlafaxine and duloxetine for neuropathic pain (Table 2)(Table 3).

The percentage of patients achieving a therapeutic dose was selected as our primary outcome because it is clinically meaningful in pursuit of efficacy. The duloxetine group had significantly more veterans achieve therapeutic dosing 120 (79.5%) versus 82/151 (54.3%) in the venlafaxine group. However, of the 120 veterans in the duloxetine group that achieved a therapeutic dose, 70 (59.3%) discontinued therapy prior to end of study with 50 (41.7%) remaining on duloxetine. Of the 82 veterans, in the venlafaxine group, that did achieve therapeutic dosing, 51 (62.2%) discontinued therapy prior to end of study with 31 (37.8%) remaining on venlafaxine therapy. Our second primary outcome was time to therapeutic dose to investigate the length of time reasonable to expect patients to wait prior to achieving the dose considered evidenced-based for their neuropathic pain. While there were several outliers in both groups remaining on subtherapeutic doses for nearly two years or more, the median time to therapeutic dose for duloxetine was 7 days with interquartile range of 44.3 days compared to median with venlafaxine 31.5 days and interquartile range of 105 days. It was somewhat surprising that both medications were titrated to therapeutic doses so quickly considering they are
antidepressants. In evaluating our primary outcomes, veterans treated with duloxetine were more likely to achieve therapeutic dose, be titrated quickly, and continue therapy all of which achieved statistical significance. This is perhaps more impressive, considering duloxetine was a restricted formulary medication requiring alternatives to be trialed before it could be approved for use. 117/151 duloxetine patients had tried venlafaxine first per formulary prior to initiating therapy with duloxetine. 35/117 (29.1%) discontinued venlafaxine after one prescription. Considering that duloxetine patients were significantly older and demonstrated sensitivity to these medications with previous failure of venlafaxine, the results clearly support duloxetine as more tolerable with a higher percentage of patients both achieving therapeutic doses and continuing therapy.

Table 2 Reasons for discontinuation and side effects

<table>
<thead>
<tr>
<th>N=302</th>
<th>Venlafaxine (151)</th>
<th>Duloxetine (151)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>D/c or expired</strong></td>
<td>N (%)</td>
<td>N (%)</td>
<td></td>
</tr>
<tr>
<td>SE</td>
<td>65 (43.1%)</td>
<td>48 (31.8%)</td>
<td>0.0569</td>
</tr>
<tr>
<td>Transferred</td>
<td>18 (11.9%)</td>
<td>9 (6.0%)</td>
<td>0.1052</td>
</tr>
<tr>
<td>Lack of efficacy</td>
<td>23 (15.2%)</td>
<td>17 (11.3%)</td>
<td>0.3962</td>
</tr>
<tr>
<td>Non-compliance</td>
<td>8 (5.3%)</td>
<td>11 (7.3%)</td>
<td>0.6366</td>
</tr>
<tr>
<td>Unknown</td>
<td>5 (3.3%)</td>
<td>14 (9.3%)</td>
<td>0.0554</td>
</tr>
<tr>
<td>Decline in renal function</td>
<td>1(0.7%)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Pain Controlled</td>
<td>0</td>
<td>2 (1.3%)</td>
<td>0.4983</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>120</td>
<td>101</td>
<td></td>
</tr>
</tbody>
</table>

Types of SE:

- Change in mood: 20 (13.3%) vs 14 (9.3%) (p = 0.3629)
- GI distress: 7 (4.6%) vs 7 (4.6%) (p = 0.7497)
- Sexual dysfunction: 6 (4.0%) vs 4 (2.7%) (p = 0.7497)
- Sedation: 4 (2.7%) vs 4 (2.7%) (p = 0.7497)
- Nightmares: 4 (2.7%) vs 2 (1.3%) (p = 0.6844)
- SI: 1 (0.7%) vs 1 (0.7%) (p = 0.7497)
- Hyperhidrosis: 2 (1.3%) vs 1 (0.7%) (p = 0.3629)
- Elevated LFTs: 0 vs 1 (0.7%) (p = 0.7497)
- Heart palpitations: 1 (0.7%) vs 1 (0.7%) (p = 0.7497)
- Allergic reaction: 3 (2.0%) vs 4 (2.7%) (p = 0.7497)
- Tremor: 2 (1.3%) vs 1 (0.7%) (p = 0.7497)
- Urinary Retention: 0 vs 3 (2.0%) (p = 0.2475)
- Myalgias: 2 (1.3%) vs 1 (0.7%) (p = 0.2475)
- Syncope: 3 (2.0%) vs 0 (p = 0.2475)
- Dizziness: 2 (1.3%) vs 1 (0.7%) (p = 0.2475)
- Hypertension: 2 (1.3%) vs 0 (p = 0.4983)
- Weight gain: 1 (0.7%) vs 1 (0.7%) (p = 0.4983)
- Insomnia: 1 (0.7%) vs 1 (0.7%) (p = 0.4983)
- Bruxism: 1 (0.7%) vs 0 (p = 0.4983)
- Headaches: 3 (2.0%) vs 0 (p = 0.4983)
- Serotonin syndrome: 0 vs 1 (0.7%) (p = 0.4983)

Table 3 Concomitant diseases and medications

<table>
<thead>
<tr>
<th>Secondary endpoints</th>
<th>Duloxetine (151 %)</th>
<th>Venlafaxine (151 %)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Comorbidities</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diabetes</td>
<td>43 (28.48)</td>
<td>27 (17.88)</td>
<td>0.0403</td>
</tr>
<tr>
<td>PTSD</td>
<td>63 (41.72)</td>
<td>70 (46.36)</td>
<td>0.4868</td>
</tr>
<tr>
<td>Depression</td>
<td>110 (72.85)</td>
<td>90 (59.60)</td>
<td>0.0206</td>
</tr>
<tr>
<td>Anxiety</td>
<td>53 (35.1)</td>
<td>43 (28.48)</td>
<td>0.266</td>
</tr>
</tbody>
</table>

**Concomitant medications**

- SSRIs: 12 (7.95) vs 17 (11.26) (p = 0.4351)
- SNRIs: 5 (3.31) vs 1 (0.66) (p = 0.2141)
- MAOIs: 0 (0) vs 0 (0) (p = 1)
- TCAs: 72 (47.68) vs 9 (5.96) (p = 0.0001)
- Gabapentin: 73 (48.34) vs 56 (37.09) (p = 0.0625)
- Pregabalin: 24 (15.89) vs 15 (9.93) (p = 0.1692)

We identified several secondary outcomes of interest including dropout rate between duloxetine and venlafaxine. Duloxetine matched its predicted rate of discontinuation with previous clinical trials with 20% failing to achieve therapeutic dosing. In contrast, venlafaxine’s rate of discontinuation was more than double that of clinical trials with 45.7% failing to achieve a therapeutic dose. Another outcome of interest was how rate of titration influenced discontinuation rate (i.e. tolerability). Duloxetine was titrated to therapeutic doses quickly (median 7 days) as expected but venlafaxine was titrated much quicker than expected (median 31 days) with 29.3% achieving therapeutic doses within 14 days and 47.6% within 30 days. The discontinuation rate with such a rapid titration was nearly 85% and did not decrease to 50% unless titrated more gradually reaching therapeutic doses after 60 days. Unfortunately, comparisons of these secondary outcomes were not sufficiently powered to evaluate significance.

There were several limitations to our study, which include its retrospective design, reliance on provider documentation for inclusion and limited external validity, as majority of patients were white males. Another limitation was due to formulary restriction, most if not all patients in the duloxetine group had to previously try and fail venlafaxine. As the patients in the duloxetine group had already not achieved benefit from a previous SNRI trial, they may be considered higher risk for a second medication failure. And furthermore, the study population was found to have a high rate of psychiatric comorbidities that if mental health was not well managed may have placed those patients at a higher risk for dropout and non-compliance.

**Conclusion**

This therapeutic impact of medication selection (TIMS) evaluation showed that in real world conditions, patients started on duloxetine for neuropathic pain were significantly more likely to achieve therapeutic dosing, be titrated to therapeutic dose much quicker, and remain on duloxetine long-term compared to venlafaxine. Based upon these results, the impact of titration schedule may play a role in discontinuation rate but requires further study to evaluate for statistical significance. We further conclude that these results indicate...
that duloxetine should be favored in treatment algorithms over
venlafaxine for neuropathic pain in formulary considerations or risk
introduction of ethical concerns into treatment.

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Conflict of interest

The authors report no conflict of interest.

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