Can we not work together to help family practitioners become more effective pain managers?

Snyder et al recently published a review in *American Family Physician* titled, “Treating Painful Diabetic Peripheral Neuropathy: An Update”, which provided an overview of pharmacologic treatment options for providers; however, some of the recommendations made by the authors were concerning. Recommendations that caught our attention included statements around pregabalin adjustment for renal impairment, using selective serotonin reuptake inhibitors (SSRIs) in the treatment of diabetic peripheral neuropathy (DPN), classification of tramadol, tapentadol, and oxycodone in DPN.

While highlighting the pharmacodynamic and pharmacokinetic differences between pregabalin and gabapentin, the statement regarding pregabalin having "no dosing adjustment requirement in patients with renal impairment" immediately caught our attention. That statement is not consistent with the current published literature and Food and Drug Administration guidance, and may therefore result in substantial confusion and adverse effects for patients if providers implement such misinformation into practice. Pregabalin is not metabolized hepatically and is excreted by the kidneys entirely unchanged and will, therefore, accumulate unless the dose is adjusted for renal dysfunction. The manufacturer labeling lists the maximum recommended dosing for pregabalin as 300 mg in two to three daily divided doses for patients with a creatinine clearance (CrCl) of 30–59 mL/min; 150 mg/day in one to two divided doses in patients with CrCl 15–29 mL/min; and 75 mg/day in patients with CrCl <15 mL/min. In patients undergoing hemodialysis, supplemental doses of pregabalin at 25–75 mg daily after hemodialysis are recommended. Renal adjustment for pregabalin is overwhelmingly recommended in the literature, and according to rates of adverse effects and concentration-dependent toxicity, failure to adjust the dose will result in accumulation and poor treatment outcomes for patients.3,4

The review article also states, “selective serotonin reuptake inhibitors and opioids are optional third-line medications”. The role of specific serotonin reuptake inhibitors (SSRIs) in treating pain associated with DPN is questionable, even to be suggested as a third-line medication. While there is a connection between mood and pain perception, SSRIs have no established mechanism or role in managing pain associated with DPN. Norepinephrine (NE) reuptake inhibition, as seen with serotonin–norepinephrine reuptake inhibitors (SNRIs) and tricyclic antidepressants, is the primary mediator of the analgesic effect. The review article highlighted that “Combined data from four small studies reveal an NNT [number needed to treat] of 7 for pain reduction with this...
class of medications”. The supporting reference is, “SSRIs have been studied in a few trials which have demonstrated a weak analgesic effect, but the clinical relevance of these compounds is questionable”, by Finnerup et al.9 There are concerns in these studies, as the methods, sample size, and results have dubious clinical relevance from which to draw conclusions about their role in DPN. The study of paroxetine for neuropathic pain involved 19 participants in an alternating crossover every 2 weeks in each arm, in which, for example, one group of six participants received a placebo for 2 weeks, followed by imipramine for 2 weeks, then paroxetine for 2 weeks.7 There are methodological weaknesses with the study, including blinding and cross-over design without a washout period, as well as utilizing a drug with long elimination half-life (19 hours), of which the therapeutic effects remained during paroxetine group’s test period. Similar to the paroxetine study, the citalopram study followed a 3-week crossover with a placebo in 15 participants.8 Finally, the study for escitalopram utilized a verbal pain rating scale (“complete”, “good”, “slight”, “none”, or “worse”) corresponding to a score of 0–11.9 Escitalopram showed improvement by 1 point with a wide standard deviation, with the largest improvements compared to placebo falling mostly in the “none” and “slight” pain relief designations. There was no statistically significant difference in quantitative sensory testing between both the groups. None of these trials utilized a parallel group with an active comparator that was selective for NE reuptake inhibition. For these reasons, SSRIs are not currently found among recommendations for any current treatment guideline for the management of neuropathic pain.10–12 As a matter of fact, the International Association for the Study of Pain cites level A/B rating for inefficacy or discrepant results for SSRIs in neuropathic pain.12 Therefore, citing SSRIs as a third-line option may be interpreted as continuing SSRIs to treat DPN in patients who are already on SSRIs for mental health indications rather than switching to evidence-based medications such as serotonin–norepinephrine reuptake inhibitors.

Another misleading statement in Snyder et al’s paper is the reference to tramadol and tapentadol as “opioid-like medications”. For clarification, tramadol and tapentadol are not “opioid-like” medications but are indeed opioid agonists in the phenylpropyl amine class.13–14 Tramadol is a μ-opioid receptor agonist with binding affinity 6,000 times lower than that of morphine and has a unique dual mechanism that includes inhibiting the reuptake of NE and serotonin.13 Tapentadol is a μ-opioid receptor agonist with binding affinity 18 times lower than that of morphine combined with NE reuptake inhibition, resulting in synergy that improves the overall potency to five times lower than oxycodone in clinical trials.15 The benefit of tramadol and tapentadol in DPN is secondary to NE reuptake inhibition, with tapentadol carrying an Food and Drug Administration-labeled indication for DPN.14–16

The authors also state, “If opioids are used, oxycodone is commonly prescribed, but alternatives include methadone, levorphanol, and morphine”. Certain opioids are presumed to be more efficacious in neuropathic pain, including those with dual mechanisms involving NE reuptake inhibition and/or N-methyl-D-aspartate.17 Methadone and levorphanol are synthetic opioids that inhibit N-methyl-D-aspartate receptor and block the reuptake of serotonin (5HT) and NE, and NE alone, respectively.18 The usefulness of oxycodone and other single mechanism opioid agonists for neuropathic pain is questionable, particularly for long-term treatment. A Cochrane review of three studies (two for DPN and one for post-herpetic neuralgia) involving 254 participants demonstrated the absence of convincing unbiased evidence supporting the benefit of oxycodone in the treatment of DPN and post-herpetic neuralgia.19

Given the shortfalls of the review by Snyder et al, we felt compelled to write a letter to the editor of American Family Practice to reconcile any misinformation and to give the authors an opportunity to refute or explain any inaccurate statements. This is particularly important since primary care physicians treat the majority of chronic pain cases in the US, with data indicating that they feel ill-prepared to do so.20,21 However, upon reviewing their submission policies, we learned that, “The first/corresponding author must be an experienced physician”. This came as a shock to all of us, as various ambiguities were glaring. Disallowing interdisciplinary participation, especially as an editorial, is an erstwhile policy in our minds and counterintuitive to the spirit of interdisciplinary medicine, which is an essential component of pain therapeutics and one of the major reasons for the opioid epidemic on our hands. An email was sent to the editors of American Family Physician explaining our concerns with a request to reconsider their policy and provide a fair peer review of our letter. We did not even receive an acknowledgement of the email after two attempts.

Ethically, we struggle with the complacence of the journal at several levels. While none of us is an MD or DO, we are well-published and have contributed as lead authors ubiquitously; we are pain clinicians, researchers, and educators, with considerable experience in training primary care physicians in pain management. We appreciate the rich
body of data indicating that interdisciplinarity results in more effective pain management. By rejecting our letter, *American Family Practice* deprived its readership of access to information from sources other than physicians who are critical to the best practice of pain medicine. Family physicians often feel isolated in their practices. Therefore, excluding nonphysicians’ input serves to perpetuate their sequestered, and not particularly effective, approaches to pain management. Finally, it appears that the editor of *American Family Practice* may have potentially attempted to obfuscate the weaknesses of the review that the journal published. If that indeed is the case, it leaves their readership with inaccurate information—and the potential to practice pain management less effectively than would be optimal. If it is not, we would welcome an editorial from their editorial staff that clarifies their intent.

In conclusion, our hope is that the editors of *American Family Practice* will not only be more diligent regarding the quality of the science published, but that journal editors and their leadership examine their collective moral compass in regard to draconian policies and potentially ensconcing the truth—as failure to do so will certainly not help family physicians gain greater competence in their efforts to manage pain, mitigate risk, and reign in a presumed opioid epidemic.

**Disclosure**

Dr. Fudin disclosed the following: DepoMed (Advisory Board, Speakers Bureau), Endo (Consultant, Speakers Bureau), Kaléo (Speakers Bureau, Advisory Board), Kashiv Pharma (Advisory Board), KemPharm (Consultant), Perrin Therapeutics (Speaker), Remitigate, LLC (Owner), and Scilex Pharmaceuticals (Consultant).

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**References**
