Comparison of duloxetine and SSRI as a treatment option of painful physical symptoms associated with major depressive disorder

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Dear editor

We would like to write about the recently published article “An observational study of duloxetine versus selective serotonin reuptake inhibitors (SSRIs) monotherapy for the treatment of painful physical symptoms in Japanese patients with major depressive disorder: primary analysis” by Kuga et al, which we read with great interest.1 The study is a good step toward finding the best treatment option for painful physical symptoms (PPSs) in patients with major depressive disorder (MDD).

Depression (MDD) is a type of mood disorder which affects individuals in such a way that they behave differently toward society. It causes abnormalities in the way of thinking, feeling, and handling daily activities of life. It is a common entity and its symptoms can be divided into two groups, namely physical (insomnia, lethargy, anorexia, PPSs) and psychological (guilt, feeling of worthlessness, agitation).2 In a cross-sectional study performed on individuals from different European countries, 50% of patients with MDD had experienced PPS as compared to normal population, where 29% of respondents reported PPS. It concluded that certain risk factors like female gender, age, and lower educational level were more likely to cause PPS.3

PPSs include pain in joints and stomach, headache, myalgia, and backache. Serotonin and norepinephrine both have a role in the pathogenesis of PPS and MDD which explains their association. There is a study4 which showed that treatment of PPS helps in control of emotional symptoms. This suggests that the patient’s capacity to achieve depression remission has a direct relation with the reduction of PPS. A treatment plan which ignores physical symptoms may result in failure of remission and can also lead to poor prognosis. Commonly used treatment options for PPS are SSRIs (fluoxetine) and serotonin-norepinephrine reuptake inhibitor (SNRI; duloxetine).

In this observational study, which claims to be the first study to compare duloxetine and SSRIs, the researchers have tried to find the best treatment option for PPS symptoms in patients with MDD. After 4 weeks, the difference in average pain score between the two groups was not substantial; however, there was an improvement in patients on duloxetine after 4–12 weeks. No serious adverse effects were reported. These results lead to the conclusion that duloxetine is promising and showed significant improvement in PPS compared to SSRIs.

If we review some previous studies which compared these two drug groups, we have different results. A study “Depression and pain: an appraisal of cost-effectiveness and cost utility of antidepressants” by Yan et al concluded that SSRIs have an edge.

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over duloxetine (an SNRI) for MDD patients with PPS keeping in view the cost utility and cost-effectiveness. Another study concluded that when PSS patients partially responding to SSRI were switched to duloxetine, there was a significant improvement in pain regardless of the switch method used.  

We suggest that similar studies are carried out in other parts of the world involving multiethnic population as results can be influenced by race/cultural differences. There is a need for further randomized studies to support the use of duloxetine over SSRIs in MDD patients with PPS.

Disclosure
The authors report no conflicts of interest in this communication.

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**Authors’ reply**

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**Dear editor**

We appreciate Drs Haider and Shafqat’s close reading of our manuscript. As they mentioned, the noradrenergic action of serotonin-norepinephrine reuptake inhibitors (SNRIs) on descending pain modulation pathways may explain its advantageous effect on painful physical symptoms (PPSs) in patients with major depressive disorder (MDD). In reference to the report by our colleagues regarding the effect of duloxetine on PPS in MDD, they stated that any treatment plan for depression that ignores physical symptoms may lead to failure of remission and poor prognosis. Harada et al in their report were very much aware of this and showed that initially the direct effect of duloxetine on PPS associated with MDD was greater than its indirect effect, whereas later the indirect effect through improvement in depressive symptoms predominated. In our study, we were concerned with showing how to optimize the treatment of MDD with PPS. This clinical question led us to plan our study comparing duloxetine and selective serotonin reuptake inhibitors (SSRIs).

After commenting on our study results, Drs Haider and Shafqat referred to two studies that appeared to have different results. First, the study by Pan et al demonstrated the superior cost-effectiveness of SSRIs over SNRIs in terms of depression and pain in Taiwan. However, it is unclear which treatment the patients in the SNRI group received, duloxetine or venlafaxine. Moreover, this was a retrospective study based on drug prescription data, and careful assessment of clinical information (ie, severity of depression and pain, dosage of each antidepressant, etc) was not in scope. Those limitations should be taken into account for appropriate interpretations.

On the other hand, the study by Perahia et al found significant improvements in PPS in MDD patients responding to SSRIs after switching to duloxetine irrespective of switching methods. This study supports usefulness of duloxetine in treating PPS as an alternative to SSRI. With that said, we would like readers to note that their study population was different from our study and their study was not designed to compare the antidepressants before and after switching. In addition, they used change in HAM-D scores as the primary objective, not pain improvement.

In order to resolve these differences, we wanted to examine pain improvement in MDD patients as the primary objective in this observational study. This is the first study, and we hope that our data may stimulate to further investigations that lead toward optimal treatment of MDD with PPS.

**Disclosure**

AK, SF, HT, AY, and RE are employees of Eli Lilly Japan K.K. TT, SH, and MM are full-time employees and stockholders of Shionogi & Co., Ltd. KT has no disclosures. TA received speakers’ honoraria from Eli Lilly Japan, K.K., Takeda Pharmaceutical Co. Ltd., Otsuka Pharmaceutical Co. Ltd., Mochida Pharmaceutical Co., Ltd., and Janssen Pharmaceutical K.K.

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