OPINION ARTICLE

Respiratory concerns of gabapentin and pregabalin: What does it mean to the pharmacovigilance systems in developing countries? [version 1; peer review: 1 approved, 1 approved with reservations]

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Abstract

Gabapentin and pregabalin, commonly known as gabapentinoids, have been widely used globally. This paper highlights the serious breathing problems due to using gabapentin and pregabalin which was warned by the United States Food and Drug Administration on December, 2019. In this article, we tried to recommend suggestions for controlling these adverse drug reactions (ADRs). Safety reports of gabapentin and pregabalin should be obtained from concerned manufacturers and reviewed for respiratory depression effects. There should be strict prescription monitoring and drug use evaluation studies. Concurrent use of gabapentin and pregabalin with other respiratory depressants should be strictly monitored. Educating patients can help in the early detection of ADRs due to gabapentin and pregabalin. Anecdotal reports on these medications should be encouraged.

Keywords

Gabapentin, Pregabalin, pharmacovigilance, Developing countries
**Introduction**

Gabapentin and pregabalin, commonly known as gabapentinoids, have been widely used globally. Gabapentin is an anticonvulsant agent used in treating various illnesses such as amyotrophic lateral sclerosis, analgesia, anxiety, neuralgia, restless legs syndrome and bipolar disorder. Pregabalin is commonly used for the treatment of painful diabetic neuropathy, fibromyalgia, diabetic neuropathy, cancer chemotherapy-induced neuropathic pain, post-herpetic neuralgia, trigeminal neuralgia, and post-operative pain. Pregabalin also acts to be an effective treatment strategy in refractory partial-onset seizures and the existing data recommends that pregabalin may be favorable as adjunctive therapy in adults with generalized or social anxiety disorder.

As per the available safety data, the use of gabapentin and pregabalin may cause neuropsychiatric related adverse drug reactions (ADRs) followed by hepatic, cutaneous and hematological reactions. Suicidal ideation, cognitive impairment, motor incoordination, dizziness are also severe forms of ADRs associated with gabapentinoids. The use of pregabalin is associated with hematological ADRs and gabapentin is also associated with liver toxicity.

**Respiratory concerns with gabapentin and pregabalin**

Respiratory depression, a highly mortal condition, due to gabapentin and pregabalin has been emerging for the past few years even in patients who were not on opioids, though post-marketing studies showed similar effects among patients taking these medications concurrently along with other respiratory suppressants. In December 2019, the United States Food and Drug Administration (US FDA) issued a drug safety alert on serious breathing problems with gabapentin and pregabalin noticed when used along with central nervous system (CNS) depressants or in patients with lung problems. US FDA reviewed data from the FDA Adverse Event Reporting System (FAERS) database of almost five years, i.e. from January 1, 2012 – October 26, 2017, which revealed 49 cases of gabapentin-induced respiratory depression. Out of 49, the majority of cases (n=34; 69.3%) were reported with pregabalin and 30.6% (n=15) cases were reported with gabapentin. Of these cases, 92% reported either a respiratory risk factor, including age-related loss of lung function, or the use of a CNS depressant. This report also revealed that 24% percent (n=12) of the 49 patients with respiratory depression died due to respiratory depression and were taking gabapentinoids.

**What do the recent respiratory problems of gabapentin and pregabalin mean for pharmacovigilance in the developing world?**

Since these ADR reports are from developed countries, it is difficult for regulators in developing countries to take decisions on these two medications that are also abundantly used in the developing world. For example, in India, there are escalating sales of gabapentin and pregabalin and while comparing the sales in 2017 to those in 2019, it was found that sales of gabapentin and pregabalin increased by 25% and 16%, respectively.

As is well documented in the literature, pharmacovigilance programs in developing countries lack robust reporting of ADRs and underreporting is a common issue. In order to bring any regulatory changes such as labeling changes or even ultimately banning the medications, one needs evidence, which largely is lacking at this point in time among developing countries.

There have often been issues like this in the past where safety concerns emerge, largely from the developed world, with drugs like selective cyclooxygenase-2 inhibitors (coxibs), cerivastatin, and glitazones, and the developing world, due to lack of stringent pharmacovigilance mechanisms, is left with little or no choice but to follow the actions taken by the developed world. In addition, developing countries lack options for communicating pharmacovigilance information among key stakeholders including consumers, which is another concern. It is astonishing that many developing countries where large quantities of medicines are used still lack strong mechanisms to monitor the safety of their products.

**Recommendations**

In the current scenario, safety reports of gabapentinoids should be obtained from concerned manufacturers and reviewed for respiratory depression effects. As understood, manufacturers are an important partner in the pharmacovigilance process with the unique advantage of formulation related information. The periodic safety update reviews (PSURs) submitted by the manufacturers of gabapentanoids can be an important source for new signal detection. In addition, the pharmaceutical manufacturer also has an obligation to report serious ADRs to the regulatory authorities.

Healthcare professionals should be watchful and report ADRs associated with gabapentinoids. Spontaneous reporting of suspected ADRs in the past have been crucial in detecting ADRs at an early stage.

Labeling changes should be made and a drug can be banned if needed. New warnings are necessary to incorporate into prescribing information including package inserts about possible respiratory depression. Concurrent use of these drugs with other respiratory depressants should be strictly monitored. Anecdotal reports on these medications should be encouraged as they can be crucial in the detection of ADRs. If noticed, causality and severity assessments should be made for the suspected ADRs.

The patients on these two medications (and the ones closely chemically related to them) should be provided with proper counseling. Educating patients can help in the early detection of ADRs and active participation of patients can help in the identification of adverse events and ADRs, description of ADRs, and ultimately prevention of drug-related harm. Hospital drug and therapeutics committees have a crucial role to play in situations like this by disseminating information within the hospital. Social media may also play a crucial role in the signal generation of suspected ADRs in these situations. Hospitals using these medications should develop risk management
plans associated with gabapentinoids usage and must disseminate any safety issues to concerned authorities.

Conclusions
Drug safety is a constantly evolving process and one must be constantly vigilant on the use of these medications. Healthcare professionals, especially prescribing physicians, nurses and pharmacists should be more cautious about using these medications in vulnerable people. Patient education and prescription restrictions of gabapentin and pregabalin may be needed until more data are available.

Data availability
No data is associated with this article.

References

8. ThePrint: US flags two top neurological drugs used in India, warns of serious breathing difficulties. 2019. Reference Source
This is an interesting article on a relevant topic. It is well written and important points are clearly presented.

Gabapentinoids, gabapentin and pregabalin, are being increasingly dispensed for indications outside US FDA labelling, particularly as part of multimodal analgesia in the perioperative setting. Emerging evidence from pre-clinical, human volunteer, inpatient, and outpatient studies suggests that their use in combination with opioids may increase the risk of respiratory depression.

In our recent studies, use of gabapentinoids on the day of surgery was associated with increased risk of postoperative pulmonary complications in patients who underwent colorectal surgery. We also found that in-hospital use of both gabapentin and pregabalin had dose dependent associations with pulmonary complications in the immediate postoperative period after total hip and knee arthroplasties.

The adverse effect of perioperative use of gabapentinoids can occur in all countries. While differences in risk across different populations are unclear (for example, the prevalence of pulmonary complications after concurrent use of opioids with gabapentinoids may differ among races), we suggest that a caution be warranted when these drugs are used in combination with opioids. Obviously, it is essential to establish effective safety reporting systems to detect signals of adverse events and disseminate the information to all relevant sections.

References

Is the topic of the opinion article discussed accurately in the context of the current literature?
Thank you for the opportunity to review this interesting short opinion paper.

The SmPCs for gabapentin and pregabalin include respiratory problems under ‘undesirable effects’; however, as the authors say, there are no recommendations to apply this to practice. This applies in all countries, particularly post-industrial societies. Our trial and observation study found too many unaddressed respiratory problems.\(^1,2\)

Whilst I concur with the sentiments expressed, I feel these should be placed in the context of full patient monitoring for the full range of possible adverse effects of prescribed medicines. Respiratory depression is extremely important, as indicated, but it is also important to check patients for other possible undesirable effects of medicines and unmedicated problems, including pain, sedation and falls. We have shown the effectiveness of this approach, and would like to see it more widely adopted\(^1,2\). Our website gives free access to our strategy.

http://www.swansea.ac.uk/adre/

References
Is the topic of the opinion article discussed accurately in the context of the current literature?
Yes

Are all factual statements correct and adequately supported by citations?
Yes

Are arguments sufficiently supported by evidence from the published literature?
Yes

Are the conclusions drawn balanced and justified on the basis of the presented arguments?
Yes

Competing Interests: No competing interests were disclosed.

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