Opioid Induced Respiratory Depression—Beyond Sleep Disordered Breathing

by Toby N. Weingarten, MD

More than a decade ago the APSF established a clear edict: “No patient should be harmed by opioid-induced respiratory depression in the postoperative period.” Research studies established a strong association between obstructive sleep apnea (OSA) and adverse postoperative opioid-related outcomes. In response, medical societies issued perioperative guidelines calling for universal screening for OSA, continuation of OSA therapies in the postoperative period, and calls for the anesthesiology team to appropriately modify the anesthetic and postoperative monitoring of patients. Unfortunately, the published rates of severe postoperative opioid-related respiratory depression (OIRD) have remained relatively constant.

More recent studies have expanded our understanding of which patients are at the highest risk for severe OIRD. These results suggest we need a more holistic approach of assessing patients beyond screening for OSA and begin to consider patient, surgical, anesthetic, and importantly anesthetic recovery characteristics. Also, these recent studies give us a better idea of when and how postoperative OIRD presents, allowing us to develop better postoperative monitoring strategies.

PATIENT CHARACTERISTICS

The association between severe OIRD and OSA is well established. For example, Mayo Clinic researchers have studied the administration of naloxone on postoperative wards as a proxy measure for severe OIRD. These studies found that patients with a history or positive screen for OSA have double the risk for developing severe postoperative OIRD compared to patients without OSA.

These Mayo Clinic naloxone studies and the PREdiction of Opioid-induced respiratory Depression In patients monitored by capnography (PRODIGY) trial have identified other important patient characteristics in addition to OSA, which also increase OIRD risk. The PRODIGY trial used bedside capnography and pulse oximetry on general care wards to identify episodes of OIRD. The PRODIGY researchers were then able to look at 46 potential patient risk factors to develop a risk score for OIRD (PRODIGY score, Table 1, next page). While, as expected, OSA and other sleep breathing disorders were found to increase risk, so was older age, male sex, congestive heart failure, and opioid-naïvety; with age beyond 70 years being most important. One weakness of PRODIGY was that many of these 46 factors were specific diagnoses and some were too rare (amyotrophic lateral sclerosis) to adequately examine. Instead, the Mayo Clinic naloxone studies used organ system disease to assess risk, and found cardiovascular disease, OSA, and debility more than doubled the OIRD risk, but that central neurologic diseases quadrupled OIRD risk. These studies suggest we should, in addition to OSA, also consider increasing age, disease burden, and debility as risk factors for OIRD.

PERIOPERATIVE COURSE

We should not just focus on patient factors when assessing OIRD risk, but also consider the perioperative course. More extensive and invasive procedures increase the risk for respiratory failure, while regional anesthetics may decrease risk. Different anesthetic drugs can increase or decrease the risk for OIRD while patients are admitted to the postanesthesia care unit (PACU). The Mayo Clinic has developed a unique protocol to manage patients in the PACU who are experiencing respiratory depression. In that protocol, OSA risk is assessed preoperatively and postoperatively. PACU nurses continuously monitor patients for episodes of respiratory depression (apnea, bradypnea, oxyhemoglobin desaturation, or “pain-sedation” mismatch (defined as when a heavily sedated patient complains of severe pain). Any patient who has one of these respiratory depressive episodes then undergoes monitoring for two additional 30-minute periods for additional episodes of respiratory depression. Those patients who have additional episodes of respiratory depression then undergo postoperative continuous monitoring with telemetry and are also considered for non-invasive positive pressure ventilation.

Figure 1. An actual capnography (1a) and pulse oximetry (1b) reading from PRODIGY, illustrates the typical OIRD breathing pattern. This patient is having repetitive apnea and partial apnea episodes interceded with normal breathing patterns. The periods of hypoxemia develop during the apnea episodes and the oxygen saturation normalizes when normal breathing resumes. Reprinted and modified with permission from Anesthesia & Analgesia and Wolters Kluwer Health, Inc.
The First Hours of Ward Admission May be Associated with Highest Frequency of OIRD

From “Respiratory Depression,”

Preceding Page

Use of the soluble volatile anesthetic isoflurane, preoperative sustained release oxycodone administration, increasing doses of intraoperative opioids, and preoperative gabapentin were all found to increase PACU respiratory depression. When one clinical area at the Mayo Clinic substituted desflurane for isoflurane and avoided routine use of midazolam, episodes of PACU respiratory depression decreased by 30%. Gabapentin and pregabalin continue to increase the risk for OIRD after PACU discharge. One study found that patients using gabapentin at home who then continued gabapentin postoperatively were at a 6-fold increase risk for naloxone administration. Researchers using the Premier Healthcare Database found that the use of preoperative gabapentin and pregabalin (as part of an Enhanced Recovery After Surgery [ERAS] multimodal protocol) increased the risk of postoperative pulmonary complications following colorectal, gynecological, and joint arthroplasty surgeries. The Federal Drug Administration has issued a black box warning that coadministration of gabapentin or pregabalin with other sedating medications increases the risk for severe respiratory complications. Given that recent meta-analyses have found that gabapentin and pregabalin are only weak analgesics when used during surgery and with evidence showing their potential to cause serious OIRD, the continued use of these medicines in ERAS protocols should be questioned.

**ANESTHESIA RECOVERY**

In many ways a patient’s course through PACU recovery can provide the most important information regarding OIRD risk on the general care wards. Patients who have PACU respiratory depression have higher rates of postoperative pulmonary complications, and as many as one third of patients who have both a positive OSA screen and PACU respiratory depression develop postoperative pulmonary complications. Further, the Mayo naloxone studies found that patients who have PACU respiratory depression have five-fold increased risk for naloxone administration. Another study which examined the postoperative course of patients administered naloxone in the PACU who then were discharged to general care wards found that these patients had a three-fold increase risk of postoperative adverse events compared to patients who did not receive naloxone in the PACU.

One possible explanation for the association between PACU respiratory depression and adverse respiratory events following discharge (even though PACU discharge criteria had been fulfilled) is that respiratory depression occurring during anesthesia recovery may persist on the ward. This was demonstrated in a study that used biowise to continuously monitor minute ventilation of 119 patients admitted to the PACU and then for the first 12 postoperative hours on the general wards. Those patients who had depressed minute ventilation in the PACU continued to do so for about 10 hours on the ward. In contrast, those patients who had normal minute ventilation in the PACU mostly continued to have normal minute volume on the wards.

**PRESENTATION OF OIRD**

Postoperative OIRD often develops in ways which are surprising to most anesthesia professionals, both as to time of onset and presenting signs and symptoms. Understanding these concepts will help develop better postoperative monitoring plans. A common belief is that critical OIRD events occur late at night when opioid analgesics, other sedating medications, and underlying OSA combine during sleep to create a lethal mix. A secondary analysis of PRODIGY found that the time relationship between OIRD, surgery, and time of day is more complex. In that study, almost all patients who had postoperative OIRD began to have multiple episodes of OIRD in the late afternoon and early evening (16:00 to 22:00) shortly after arriving on the wards. The frequency of OIRD episodes surged during the early morning hours (02:00 to 06:00). However, in the Mayo Clinic naloxone studies, naloxone was typically administered during the afternoon and evening. These studies suggest it is the first few hours of admission to the ward that are most hazardous. Therefore, monitoring for OIRD should begin upon admission to the ward and not wait until bedtime.

Another common belief is that OIRD usually presents as bradypnea and/or hypoxemia. However, studies which examined nursing notes preceding severe episodes of OIRD have found that oftentimes normal respiratory rates and oxygen saturations are documented. There are several potential explanations for these findings. One is that severe OIRD develops suddenly, and thus, signs of respiratory depression are not present during preceding vital signs checks. Research does not support this possibility. Postoperative OIRD persists for hours following PACU discharge, and PRODIGY showed that patients usually have multiple, repetitive OIRD events. A more likely possibility is that bradypnea or oxygen desaturation is a different picture of OIRD than is com-

### Table 1: PRODIGY Scoring System for Assessing Risk for OIRD Among Patients Hospitalized on General Care Wards Receiving Opioids

<table>
<thead>
<tr>
<th>Clinical characteristic</th>
<th>PRODIGY score†</th>
<th>RD risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low risk</td>
<td>&lt; 8</td>
<td>REF</td>
</tr>
<tr>
<td>Intermediate risk</td>
<td>8–14</td>
<td>2-fold</td>
</tr>
<tr>
<td>High risk</td>
<td>≥ 15</td>
<td>6-fold</td>
</tr>
</tbody>
</table>

**PRODIGY category**

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<td>High risk</td>
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</tbody>
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**Abbreviations:** PRODIGY, Prediction of Opioid-induced respiratory Depression in patients monitored by capnoGraphy; RD, respiratory depression; REF, reference range.

*Sleep-disordered breathing can be determined from either patient history or positive screen for sleep apnea.

To calculate the PRODIGY risk score, summate the assigned points per positive clinical characteristic. Patients are assigned low-, intermediate-, or high-risk category based on the number of points. Compared to low-risk scored patients, intermediate-risk patients have a 2-fold increase and high-risk patients a 6-fold increased risk for experiencing respiratory depressive episodes on the general care ward. (Adapted from Khanna et al.)

See “Respiratory Depression,” Next Page
From “Respiratory Depression,” Preceding Page

From “Respiratory Depression,” Preceding Page

monly assumed. In PRODIGY, almost 100% of OIRD episodes consisted, in part, of an apnea or partial apnea event, and isolated bradypnea or oxygen desaturation were extremely rare (Figure 1). Though not shown, patients who were on supplemental oxygen and had OIRD, oftentimes did not have periods of oxygen desaturation during apnea spells. In the setting of a repetitive apnea OIRD breathing pattern, it is plausible that when a nurse comes to make an assessment, the patient will awaken to the point that normal breathing resumes, thus masking signs of respiratory depression. It is important to note that in many cases of severe OIRD, the nursing notes, while not recording signs of respiratory depression, will note that a patient is somnolent or sedated. These observations suggest that nurses should be trained to quietly observe breathing patterns of sleeping patients to assess respiratory status before measuring other vital signs that may awaken the patient such as blood pressure measurement. The fact that many patients who developed critical OIRD events were noted to be somnolent or sedated beforehand also presents an opportunity to educate nursing staff that such sedated patients should be considered higher risk and more carefully monitored.

A PROPOSED NEW APPROACH TO POSTOPERATIVE OIRD

Findings from these recent studies can allow the anesthesia professional to expand the assessment of OIRD risk beyond a preoperative OSA screen (Figure 2). In addition to a mandatory preoperative screening of patients for OSA, risk for OIRD should consider advancing age and overall disease burden. Calculating the PRODIGY score for OIRD risk is easy, convenient, and can be incorporated into electronic health record platforms. Patients with OSA should continue to use their continuous positive airway pressure or other devices in the postoperative period. The anesthetic could be modified for higher risk patients utilizing regional blocks, shorter acting agents, and nonsedating analgesics (e.g., acetaminophen). Finally, during anesthesia recovery, patients should be monitored for episodes of respiratory depression. Based on this information as well as the extent of the surgical procedure, the anesthesia professional could tailor the postoperative care plan based on level of risk in regards to postoperative disposition and level of monitoring where patients deemed higher risk for OIRD are specifically targets for escalation of postoperative care.

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