Problem

Neuromuscular blocking agents are high-alert medications because of their well-documented history of causing catastrophic injuries or death when used in error. These drugs are used during tracheal intubation, during surgery of intubated patients, and to facilitate mechanical ventilation of critically ill patients. However, neuromuscular blockers have been inadvertently administered to both adult and pediatric patients who were not receiving proper ventilatory assistance. Because neuromuscular blockers paralyze the muscles that are necessary for breathing, some patients have died or sustained serious, permanent injuries if the paralysis was not witnessed by a practitioner who could intervene.

After a patient receives a neuromuscular blocker, progressive paralysis develops, initially affecting the small muscle groups such as the face and hands, then moving to larger muscle groups in the extremities and torso until all muscle groups are paralyzed and respiration ceases. However, full consciousness remains intact, and patients can experience intense fear when they can no longer breathe. They can also sense pain. The experience can be horrific for patients and can lead to psychological trauma, including post-traumatic stress disorder.1

The ISMP National Medication Errors Reporting Program (ISMP MERP) has received well over 100 reports of errors involving neuromuscular blockers. But the true incidence of injuries from erroneous administration of neuromuscular blockers is much higher than reflected in our error-reporting program. Although some errors have occurred during anesthesia in the operating room (OR), many have taken place outside this setting, in emergency departments (EDs), interventional radiology departments, intensive care units (ICUs), and other medical, surgical, and psychiatric units.

The most common type of error with neuromuscular blockers appears to be administration of the wrong drug. A 2009 analysis of 154 events over a five-year period showed that neuromuscular blockers were not the intended drug in approximately half of all wrong-drug errors.2 Practitioners thought they were administering a different drug, so patients may not have been supported with mechanical ventilation. More than 80% of these wrong-drug errors reached the patient, and approximately one-quarter resulted in patient harm—a significantly high rate when compared to the fewer than 1% of events causing harm with all other wrong-drug errors during the same study period.2

Errors with neuromuscular blockers can be attributed to one or more common causes. The following is a sample of the causes of errors with examples:

- **Look-alike Packaging and Labeling**
  An ED nurse administered pancuronium instead of influenza vaccine to several patients. The vials were the same size and the labels were quite similar. The look-alike vials had been stored next to each other in the refrigerator. The patients experienced dyspnea and respiratory depression but, fortunately, they sustained no permanent injuries.

- **Unsafe Mnemonics**
  A ventilated ICU patient was receiving vecuronium and a potassium chloride infusion. After the patient was extubated, an infusion bag containing vecuronium remained in the room and was mistaken as a potassium chloride infusion. Soon after the medication was started, the patient arrested, requiring intubation and ventilation for six more hours.

- **Drug Administration After Extubation**
  An anesthesiologist was interrupted while preparing syringes of midazolam and rocuronium. Anesthesiology computer rule after entering “cis” completed the drug field name with cisatracurium. This generated a label for the neuromuscular blocker, which was prepared and dispensed.

- **Unlabeled and Mislabeled Syringes**
  Prefilled syringes of saline flushes were not available in the ED, so nurses prepared a supply each day from multiple-dose vials. Vecuronium had recently been prepared for a trauma patient in the ED, but it was not used. The syringe was not labeled and was inadvertently placed with the saline flush syringes. The syringe containing vecuronium was later used to flush the intravenous (IV) line of a 3-year-old child. The child became flaccid and stopped breathing. She was quickly intubated and ventilated, so permanent harm was averted.

- **Unlabeled and Mislabeled Syringes**
  Anesthesiology was interrupted while preparing syringes of midazolam and rocuronium.2 When he returned, he...
administered the contents of one syringe to a patient in the holding area, believing it contained midazolam. He was again called away, and when he returned, the patient was unresponsive. The patient was intubated and given a reversal agent, and surgery was postponed. It was later determined that the anesthesiologist had administered the syringe containing succinylcholine.

**Unsafe Storage**
Atracurium was administered instead of hepatitis B vaccine to several infants, who developed respiratory distress. One infant sustained permanent injury and another infant died. Neuromuscular blockers had never been available as unit stock in the nursery. An anesthesiologist from a nearby OR had placed the atracurium vial in the nursery refrigerator near look-alike vaccine vials. Similar mix-ups with vaccines continue to occur.1

**Orders Entered Into Wrong Electronic Health Records**
A medical resident electronically prescribed vecuronium for the wrong patient with a similar name, who was located on a medical unit. The correct patient was ventilated and in the ICU. The pharmacist and technician did not question the infusion for a medical unit patient. An independent double-check was carried out by two nurses before administration, but neither nurse was aware that the patient required ventilation with this drug.

**Knowledge Deficit on Drug Action And Required Ventilation**
An ED physician gave a verbal order for a trauma patient to receive vecuronium and midazolam, which were administered prior to intubation. He then mistakenly entered electronic orders for these medications into another patient’s record. An ED nurse administered the medications to the patient without recognizing that vecuronium would paralyze the respiratory muscles. After she left the room, the patient arrested. The ED team responded, but the patient could not be resuscitated.

**Syringe Swaps**
Succinylcholine was inadvertently administered instead of fentanyl prior to the induction of anesthesia.2 The anesthesiologist had drawn both drugs into 2-mL syringes, and had applied a blank red-

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**Primary Recommendations**

**Limit Access**
Eliminate the storage of neuromuscular blockers in areas of the hospital where they are not needed.2 Allow unit stock only in the OR, ED, and critical care units where patients can be properly ventilated and monitored. Consider limiting the number of neuromuscular blockers on formulary, and eliminate storage from pharmacy stock when possible. Regularly review these storage areas, both inside and outside of the pharmacy, including agents that require refrigeration, and consider the potential for mix-ups. Limiting access to these products is a strong deterrent to inadvertent use.

**Standardize Prescribing**
Outside the OR or procedural areas, orders for neuromuscular blockers should only be part of an intubation protocol, or an order set to maintain a specific level of paralysis while the patient is on a ventilator only. Do not accept neuromuscular blocker orders for “use as needed for agitation.” Include the need for ventilation support during and after administration and automatic discontinuation of these agents in electronic records after extubation and removal from a ventilator. Completely disallow orders to “resume the same medications” upon patient transfer.

**Build Computer Alerts**
Build alerts in the computer system to verify the patient’s location when neuromuscular blocker orders are being prescribed or entered/verified by pharmacy. If the patient is not in a critical care unit, ED, OR, or invasive procedure area, prescribers should verify that they are entering the order into the correct patient profile, and pharmacists should question the order and verify ventilatory assistance before dispensing the drug. If possible, establish computerized cross-checking of the patient’s location when entering neuromuscular blocker orders (as with
other drugs limited to administration on a specific unit). Cautionary messages may also appear on automated dispensing cabinet (ADC) screens.

**Segregate Storage**
Segregate, sequester, and differentiate all neuromuscular blockers from other medications, wherever they are stored in the organization. In areas where they are needed, place neuromuscular blockers in a lidded box or in a rapid sequence intubation (RSI) kit; one option is a highly visible red-orange storage container available commercially. If neuromuscular blockers must be stored in ADCs, keep them in separate lidded pockets, away from other drugs. Also segregate neuromuscular blockers from all other medications in the pharmacy by placing them in separate lidded containers in the refrigerator or another secure, isolated storage area. Organize anesthesia carts and trays to avoid the proximity of look-alike vials, syringes, or bags, and display the labels so they are readily visible.

**Assess Labeling and Packaging**
Require a medication safety officer (MSO) and an anesthesia staff member to evaluate any new neuromuscular blocker’s packaging and labeling prior to procurement, and introduce auxiliary label enhancements and education, if necessary, before distribution. Use brands of neuromuscular blockers that clearly differentiate the vials from other products via warnings on the label, vial cap, and metal ferrule around the rubber stopper. (All manufacturers of these agents are required to provide cautionary labeling. The development of a universal symbol for neuromuscular blockers remains to be determined.)

**Affix Warning Labels**
Place auxiliary labels on all storage bins and final medication containers (e.g., vials, syringes, IV bags) of neuromuscular blockers that state “WARNING: PARALYZING AGENT—CAUSES RESPIRATORY ARREST” to clearly communicate that respiratory paralysis will occur and ventilation is required. The warning labels should not cover important label information. For infusions, one hospital system also places a warning on a port tag that will be seen by nurses when they spike the bag to attach tubing. The use of shrink-wrap sleeves is questionable because they make all vials look alike.

**Dispense From Pharmacy**
For nonurgent doses in the OR or ED, and continuous infusions in the ICU, dispense neuromuscular blockers from the pharmacy in the most ready-to-use form. The Anesthesia Patient Safety Foundation recommends the use of labeled, prefilled syringes and prepared infusions of neuromuscular blockers (and other anesthesia drugs) dispensed by pharmacy, commercially available, or outsourced, rather than self-prepared syringes or infusions. Properly labeled, prefilled syringes have the potential to improve system safety, reduce syringe swaps, and enhance work efficiency. Never dispense a neuromuscular blocker to a unit that cannot support mechanical ventilation.

**Verify Neuromuscular Blockers**
Remind practitioners that reading labels is the first defense for avoiding errors. Equally important, given human fallibility, is the implementation of point-of-care barcode scanning to verify neuromuscular blockers and patients before administration. In the OR and procedural areas, if barcode scanning is not undertaken, consider alternative verification systems including speakers and touch screens that provide automatic auditory and visual verification of drugs and important alerts prior to administration.

**Use Smart Infusion Pumps**
Administer all neuromuscular blocker infusions via a programmable smart infusion pump using dose error-reduction software. Smart infusion pumps should be programmed to allow the selection of a neuromuscular blocker infusion only in patient-care areas that are capable of caring for ventilated patients receiving such agents. When a neuromuscular blocker is selected in units where ventilation is possible, a clinical advisory warning should note that the drug paralyzes the respiratory muscles, and the nurse must confirm that the patient is on mechanical ventilation. The flow rate of infusions of neuromuscular blockers should be presented, and entered into the pump using the same standard dosing units that prescribers use (e.g., mcg/kg/minute vs. mcg/kg/hour).

**Use Clear Terminology**
Always refer to these drugs as “neuromuscular blockers” or “paralyzing agents.” Never call them “muscle relaxants.”

**Secondary Recommendations**

**Reduce Unsafe Mnemonics**
Review order-entry systems to identify problematic mnemonic auto-fill entries and label generation associated with neuromuscular blockers, and implement safer computer rules for mnemonics when indicated.

**Provide Warnings on Pharmacy Labels**
Ensure that pharmacy work labels and infusion/product labels for neuromuscular blockers are clear and accurate, and contain all necessary warnings.

**Require Proper Labeling**
Promote accurate labeling of all infusions and syringes containing neuromuscular blockers both in the OR and in patient-care locations outside the OR. (When possible, prepared and labeled syringes and bags should be provided.)

**Provide Access to Reversal Agents**
Ensure that all appropriate reversal agents for neuromuscular blockade are available to qualified staff who might need them in an emergency. In protocols, identify who is permitted to administer the reversal agent in an emergency and provide readily available instructions for administration.

**Flush the Line**
If a neuromuscular blocker has been administered, all of the drug should be flushed from the IV line or the line should be changed (and any source container removed) prior to extubation.

**Timely Dispensing and Prompt Removal**
Pharmacy should practice just-in-time dispensing of neuromuscular products when possible to avoid unnecessary access to these products before use. When the drugs are no longer needed, place unused/partially used vials, bags, and syringes of neuromuscular blockers in a sequestered bin for return to the pharmacy. Unused patient-specific doses should be destroyed/discarded after the

continued on page 107

Vol. 44 No. 3 • March 2019 • P&T® 93
patient has been extubated or the drug has been discontinued.

**Increase Awareness**

Educate staff about the risk of serious errors with these high-alert drugs. Provide staff with a list of both generic and brand names for all neuromuscular blockers available at your location, and include usual dosages and any special guidelines associated with their preparation, distribution, administration, and monitoring. In addition, use the information provided here to assess your safety practices.

**Verify Competency**

Establish a formal training program and competency verification process for practitioners who are involved in preparing, dispensing, and administering neuromuscular blockers. These drugs should only be administered by staff who have experience in maintaining an adequate airway and respiratory support, and only in units where intubation and respiratory support can be provided.

**REFERENCES**

8. In 2019, ISMP is celebrating its 25th anniversary of helping health care practitioners keep patients safe and leading efforts to improve the medication-use process. For more information, visit www.ismp.org.