Using a Patient Safety Analysis to Guide Infusion Therapy for Patients With COVID-19

Mary Hagle, PhD, RN, NPD-BC, FAAN • Kate Snyder, PharmD, BCPs • Karen Janicek, BSN, RN • Kimberly Bell, PharmD • Beth Dietz, MSN, RN, CCNS, CCRN-K • Nathan Gundacker, MD • Angelina Kinter, MSN, RN, CCRN-K • Annette Severson, EdD, RN

ABSTRACT

In the United States, during the Coronavirus Disease 2019 (COVID-19) pandemic, patients with COVID-19 overwhelmed available intensive care beds, staffing levels were unpredictable, and personal protective equipment was limited. The safety of situating electronic infusion pumps outside patient rooms was evaluated using an internal risk assessment. Based on a low level of risk, a procedure was developed to direct clinicians as to when this process is appropriate during a national crisis. A standardized analysis, Healthcare Failure Mode and Effects Analysis, was conducted to identify all potential risks and implement actions that would eliminate or control the risk. No adverse events were reported. Safe systems and preparation can protect patients.

Key words: COVID-19, HFMEA, ICU, infusion, patient safety, personal protective equipment, PPE, pump
DURING NATIONAL AND WORLD HEALTH CRISSES, SUCH AS THE CORONAVIRUS DISEASE 2019 (COVID-19) PANDEMIC, EMERGENCY MEASURES IN HEALTH CARE PROCESSES ARE OFTEN NECESSARY. IN THE UNITED STATES IN 2020, COVID-19 PATIENTS OVERWHELMED AVAILABLE INTENSIVE CARE UNIT (ICU) BEDS; STAFFING LEVELS WERE UNPREDICTABLE; AND PERSONAL PROTECTIVE EQUIPMENT (PPE), CRITICAL MEDICATIONS, AND SUPPLIES WERE LIMITED. THE SITUATION WAS SO DIRE THAT THERE WAS A NATIONAL CALL FOR IDEAS TO CONSERVE PPE FROM THE EDITORS OF THE JOURNAL OF THE AMERICAN MEDICAL ASSOCIATION ON MARCH 20, 2020.1 ADDITIONALLY, INDUSTRY DOCUMENTS, ANECDOTAL DESCRIPTIONS ON SOCIAL MEDIA, AND PROFESSIONAL WEBSITES PROVIDED RESOURCES AND NOVEL PRACTICES TO CARE FOR PATIENTS WITH COVID-19. THE NATIONAL COVID-19 BED CAPACITY, CLINICAL STAFF SHORTAGE, AND MEDICATION/SUPPLY SHORTAGE WERE ECHOED AT THE AUTHORS’ MIDWESTERN ACADEMIC MEDICAL CENTER.

BECAUSE THE SARS-COV-2 VIRUS, WHICH CAUSES THE COVID-19 DISEASE, CAN BE AIRBORNE, CRITICALLY ILL PATIENTS WITH COVID-19 WERE PLACED IN NEGATIVE PRESSURE ROOMS, FORMALLY KNOWN AS AIRBORNE INFECTION ISOLATION ROOMS. THESE ROOMS CONTAIN CONTAMINATED AIR FROM THE PATIENT THAT IS FILTERED BEFORE IT IS BLOWN OUTSIDE. THE DOOR MUST BE KEPT CLOSED TO MAINTAIN THE NEGATIVE AIR PRESSURE. WITH ALL EXISTING NEGATIVE PRESSURE ROOMS IN USE, ADDITIONAL ROOMS WERE CONVERTED TO NEGATIVE AIR PRESSURE USING FREE-STANDING IMPROVISED AIR FILTRATION AND FAN SYSTEMS. HOWEVER, SEVERAL CHALLENGES WERE NOTED. FIRST, THE FAN SYSTEMS WERE OFTEN NOISY. SECOND, WITH ROOM DOORS CONSTANTLY CLOSED, ALARMS WERE DIFFICULT TO HEAR OUTSIDE OF THE ROOM.

STRICT ISOLATION PROTOCOLS WERE IN PLACE FOR STAFF SAFETY. CLINICIANS WORKING IN COVID-19 ICU ROOMS WERE OBSERVED WHILE THEY DONNED APPROPRIATE PPE TO ENSURE THAT THEY WERE PROTECTED WHEN THEY ENTERED A PATIENT ROOM AND AGAIN WHILE THEY DOFFED PPE AFTER COMING OUT OF THE ROOM. THIS PROCESS TOOK TIME BUT WAS CRITICAL IN PROTECTING STAFF. OFTEN, NURSES REMAINED IN THE ICU ROOMS FOR 4 OR MORE HOURS AT A TIME BECAUSE OF THE PPE SHORTAGE AND TIME TO DON AND DOFF. OTHER ISSUES AROSE WHILE WORKING IN THESE ROOMS. CLINICIANS BECAME VERY WARM WHILE THEY WORKED, FACE MASKS OR SHIELDS CHAFED THEIR SKIN, AND THE POWERED AIR-PURIFYING RESPIRATORS DRIED THEIR EYES. FREQUENT MEDICATION AND/OR RATE CHANGES ALSO KEPT NURSES IN THE ROOM FOR EXTENDED PERIODS OF TIME.

SEVERAL CLINICIANS SAW JOURNAL COMMENTS AND POSTS ON SOCIAL MEDIA ABOUT NURSING STAFF SITUATING ELECTRONIC INFUSION PUMPS FOR INTRAVENOUS (IV) THERAPIES OUTSIDE COVID-19 ICU PATIENT ROOMS. IN THE INTEREST OF CONSERVING PPE SUPPLIES AND INCREASING STAFF EFFICIENCY, QUESTIONS WERE RAISED AS TO WHETHER THIS STRATEGY COULD SAFELY BE ADOPTED AT THE AUTHORS’ MEDICAL CENTER. ALTHOUGH THE IMMEDIATE BENEFITS APPEARED TO BE CLEAR, A TASKFORCE WAS FORMED TO VALIDATE THE FEASIBILITY AND SAFETY OF THIS NEW PRACTICE AND THEN DEVELOP A PROCEDURE IF INDICATED. AN INTERNAL HEALTH AND SAFETY RISK ASSESSMENT WAS COMPILED WITH RELATED RISK CONTROL MEASURES. A FORMAL RISK ANALYSIS, KNOWN AS A HEALTHCARE FAILURE MODE AND EFFECTS ANALYSIS (HFMEA), WAS ALSO CONDUCTED. ELECTRONIC INFUSION PUMPS WERE APPROVED TO BE SITUATED OUTSIDE THE COVID-19 ICU PATIENT ROOMS, AND A PROCEDURE WAS DEVELOPED FOR USE DURING THE NATIONAL CRISIS. THE PURPOSE OF THIS ARTICLE IS TO DESCRIBE THE PROCEDURE, REPORT THE FINDINGS OF AN HFMEA, AND SHARE OUTCOMES OF THIS PROCEDURAL CHANGE.

SITUATING INFUSION PUMPS OUTSIDE COVID-19 ICU PATIENT ROOMS

IN RESPONSE TO QUESTIONS ABOUT SITUATING ELECTRONIC INFUSION PUMPS OUTSIDE COVID-19 ICU PATIENT ROOMS, AN INTERDISCIPLINARY TASKFORCE WAS CONVENED TO EXAMINE RISKS AND PROCESSES OF THIS PROCEDURE. THE TASKFORCE WAS LED BY A NURSE INFECTION PREVENTIONIST AND CONSISTED OF AN ICU CLINICAL NURSE SPECIALIST, MANAGERS, NURSE SCIENTIST, PHARMACISTS, AN INFECTIOUS DISEASE PHYSICIAN, AND OTHER ANCILLARY STAFF. THE 10 TASKFORCE MEMBERS REPRESENTED THE FOLLOWING DEPARTMENTS: BIOMEDICAL ENGINEERING, FACILITY MANAGEMENT, INDUSTRIAL HYGIENE, INFECTIOUS DISEASE, NURSING, PHARMACY, AND QUALITY MANAGEMENT (WHICH INCLUDES INFECTION CONTROL, MEDICATION MANAGEMENT, AND PATIENT SAFETY).

SEVERAL VIRTUAL AND FACE-TO-FACE MEETINGS WERE HELD OVER THE COURSE OF 3 WEEKS TO IDENTIFY KNOWN OR FORESEEABLE HAZARDS AND TO DESCRIBE THESE RISKS USING AN INTERNAL HEALTH AND SAFETY RISK ASSESSMENT TOOL. GLOBAL QUESTIONS INCLUDED: “WHAT COULD HAPPEN?” “HOW COULD IT HAPPEN?” AND “ARE THERE ANY PARTICULAR FACTORS OR ISSUES THAT CONTRIBUTE TO THE RISK(S)?” DURING THE MEETINGS AND VIA EMAIL, TASKFORCE MEMBERS CONSIDERED POSSIBLE ADVERSE OUTCOMES THAT COULD OCCUR FROM SITUATING INFUSION PUMPS OUTSIDE COVID-19 ICU PATIENT ROOMS. POTENTIAL ADVERSE OUTCOMES INVOLVED EVENTS RELATED TO, FOR EXAMPLE, TRIP HAZARD, TUBING DISCONNECTION, INFECTION, DELAY OF MEDICATION ADMINISTRATION, INACCURATE INFUSION RATES, MEDICATION WASTE, AND DEPLETION OF IV EXTENSION TUBING. IT WAS NOTED BY SOME TASKFORCE MEMBERS THAT THE PRACTICE OF RUNNING IV EXTENSION TUBING UNDER A DOOR HAD BEEN USED INFREQUENTLY BUT SAFELY FOR PATIENT CARE DURING MAGNETIC RESONANCE IMAGING SERVICES.

DISCUSSION WITH FACILITIES MANAGEMENT STAFF HELPED THE TASKFORCE DECIDE THAT HAVING IV EXTENSION TUBING RUN UNDER THE DOOR WOULD HAVE NO IMPACT ON NEGATIVE AIR PRESSURE.

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Corresponding Author: Mary Hagle, PhD, RN, Clement J. Zablocki VA Medical Center, Nursing Education & Research, 5000 W National Ave, Milwaukee, WI 53295 (mary.hagle@va.gov).

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After deliberating, the taskforce agreed that the likelihood of an adverse event occurring was minimal and the consequences should an event occur would likely be minor. Furthermore, the taskforce brainstormed actions to eliminate or control other risks. Logistics ensured that a 2-month supply of extension tubing would be on hand, and pharmacy identified medications with critical shortages that would not be infused outside the room. Based on the summary of the risk assessment and the risk control measures, the procedure of situating infusion pumps outside COVID-19 ICU patient rooms was supported by clinical and administrative leadership.

Following leadership endorsement, a procedure was swiftly developed and promptly initiated. It outlined the process for implementation in patient rooms and identified risk control measures with available evidence, including alerts from the ECRI Institute, Infusion Nurses Society, and the Institute for Safe Medication Practices. The procedure was indicated only for COVID-19 ICU patient rooms (Table 1).

Specifically, the procedure contained an initial checklist of applicable needs or conditions to situate the infusion pump outside the COVID-19 ICU patient room. If the decision was made to situate the infusion pump outside the room, the procedure stated that the patient must have a central venous catheter. The remaining procedure document was grouped into 3 topics: IV tubing considerations, infusion pump considerations, and medication directions. For example, regarding medication directions, infusing propofol outside the room generated much discussion because of safety concerns and a medication shortage at that time; it was decided not to infuse propofol outside the room.

The procedure was not a plan for total patient care: it did not address expected nursing practices such as appropriate and timely patient assessments, cautions when administering a medication the first time, and actions to take for patient responses to medications or procedures. The procedure was revised slightly after 1 month for readability and to acknowledge a new patient safety notice from the Department of Veterans Affairs to all patient safety managers. This safety notice warned that situating infusion pumps outside of patient rooms should only be used as a last resort to conserve PPE. The conservation of PPE was already a condition of implementation; thus, no other changes were needed.

**HFMEA FOR PATIENT SAFETY ANALYSIS**

Soon after approval of the practice, nurses in the COVID-19 ICU successfully situated 1 pump outside a patient’s room without any adverse events. To consider all potential vulnerabilities and gaps associated with situating infusion pumps outside COVID-19 ICU patient rooms, the patient safety managers recommended completing an HFMEA. Although this analysis is best performed before starting a new practice, the taskforce supported conducting this standardized patient safety analysis even after implementation of the procedure because there was minimal evidence to support this practice and benefits seemed to outweigh the risk.

**Description of an HFMEA**

An HFMEA is a tool and an approach that is often utilized to proactively assess vulnerabilities and identify potential problems in health care, particularly those relating to patient safety, before they occur and reach the patient. The Failure Mode and Effects Analysis (FMEA) model has been used within the engineering community for >30 years. Within the Department of Veterans Affairs, the National Center for Patient Safety adapted the FMEA model specifically for health care, resulting in the HFMEA acronym.

An HFMEA for a new process is composed of a series of activities that are systematically conducted to arrive at actions for mitigating risks. Although different numbers of actions have been reported, from 5 steps to 9 phases, the activities of HFMEA are similar. Irrespective of how many steps or phases, all HFMEA activities are needed to complete the hazard or risk analysis and identify appropriate actions to be taken.

**Steps of an HFMEA**

In brief, the following 5 steps represent the HFMEA that was conducted by the authors:

1. Define the HFMEA topic
2. Assemble the team
3. Graphically describe the process
4. Conduct the hazard analysis
5. Determine actions and outcome measures.

One of the authors (K.S.) was a patient safety fellow at the time and initiated the HFMEA on this procedure change for taskforce review. First, the topic was defined as situating electronic infusion pumps outside of COVID-19 ICU patient rooms with long IV extension sets and/or multiple IV extension sets. This procedure was done in an effort to conserve PPE during the COVID-19 pandemic by minimizing entry into the patient room and to decrease caregiver exposure by reducing the amount of time in patient rooms.

Second, the HFMEA pump team was composed of the original taskforce members with additional experts from the medical center. In step 3, starting with the written procedure, a detailed description of the process was outlined. The procedure within the HFMEA was separated into 5 sections, starting with a decision to situate the infusion pump outside the room and ending with preparation to infuse medications and follow-up actions (Table 2).

In step 4 of the HFMEA, a hazard analysis was conducted; this consists of 2 parts. First, potential failure modes or hazards need to be identified; second, the failure mode is evaluated to determine the likelihood of its occurrence.
**TABLE 1**

Procedure for Electronic Infusion Pumps Outside COVID-19 ICU Patient Rooms

This procedure is only for the COVID-19 ICU until further notice. Initial approval of procedure: 4/14/20; Revised: 5/19/20

I. Patient considerations for having an IV pump outside an ICU COVID-19 room:

- **PUMP OUTSIDE ROOM:**
  - Need to conserve PPE
  - Multiple medications with frequent changes
  - Patient requiring care from more than 1 nurse
  - Sufficient administration extension sets for all meds

- **NOTE:** There is a Patient Safety Notice: this practice is not endorsed by Veterans Health Administration National Infectious Diseases Service. "Use of infusion pumps outside of patient rooms should only be used as a last resort, after all other efforts to conserve PPE have been exhausted, and in the very specific incidence of critical facility level PPE shortages."

- **PUMP INSIDE ROOM:**
  - Sufficient PPE or
  - Patient care and IV med/pump monitoring can be bundled

II. IV Pump outside an ICU COVID-19 room implementation:

A. Patient must have a CVC

B. IV tubing considerations

1. Use microbore tubing if possible, although large bore can be used
   - Using 20 feet of microbore tubing: accuracy was within specifications when infusion rates were between 5 and 300 mL/h
     - Less than 5 mL/h: infusion slow and occlusion alarm may be delayed
     - More than 300 mL/h: higher back pressure so more frequent alarms

2. Use triple port connector if needed
3. Include connector volume for priming volume of extension sets
4. Label tubing inside/outside room
5. Ports: use disinfection caps on all ports
6. Ports: do not use any y-sites or needle-free access ports that have been on floor.
7. Change IV administration sets and intermittent tubing every 7 days (along with the CVC dressing) to minimize handling.
8. Secure tubing to prevent disconnection or trip hazard:
   - Cover tubing on floor with orange cord protector from operating room (trip cord protector, 8 × 24 inches)
   - Post signage
9. Develop disinfection/cleaning schedule for floor around tubing/protector.
   - Unknown: absorption from cleaning agents: ECRI recommends clean around tubing.

C. IV pump considerations

1. Occlusion pressure settings: follow manufacturer’s recommendation to accommodate adjusting occlusion pressure rates based on resistance, low infusion rates, high infusion rates through CVC.
2. Patient identification (wristband) on patient and pump
3. Utilize designated pumps that are conducive for extension tubing; do not use syringe type pump set
4. No need to lock pump if visitors and patients are not walking around
5. Some hospitals keeping controlled substance infusions in room
6. Document in chart that IV pump is outside room

D. Medication directions

1. Prime long extension sets with medication before connecting to patient to minimize risk of delayed start
2. Flush rate to be the same as medication delivery rate so the medication infuses over ordered duration.
3. Medications that CANNOT be infused from the extended infusion set: propofol (because of shortage), blood, fat emollients, others as identified; insulin infusion

4. Compatibilities:
   - Fentanyl is y-site compatible with dexmedetomidine and midazolam
   - Ketamine is y-site compatible with midazolam
   - Dexmedetomidine is y-site compatible with fentanyl and midazolam
   - Midazolam is y-site compatible with dexmedetomidine, fentanyl, and ketamine

III. Alternatives to IV pumps:

- If using gravity drip: must count drops, even with flow regulator device; rate may be slower because length of tubing: recalculate
- Hypodermoclysis for hydration

Abbreviations: CVC, central venous catheter; ICU, intensive care unit; IV, intravenous; PPE, personal protective equipment.

Resources and information from: ECRI, Infusion Nurses Society, Institute for Safe Medication Practices.

**NOTE:** Procedure developed by staff from the following departments: nursing, pharmacy, quality management and safety, infectious disease, biomedical engineering, facility management, and industrial hygiene. The contents do not represent the views of the US Department of Veterans Affairs or the US government.

and the severity of injury if an injury occurs. Thus, in part 1, potential failure modes were identified for each of the 5 sections of the procedure (Table 3). For example, in section 3, preparing and using the infusion pump, there were 3 potential failure modes: staff are unfamiliar with infusion pump manufacturer recommendations; wristbands are not attached to patient and/or not taped to pump; and staff are not familiar with different pump types.
| Step 3: Describe the process in detail: based on the procedure for situating IV infusion pumps outside COVID-19 ICU patient rooms |
|---|
| **1. Deciding if infusion pump can be outside patient room** |
| A. Determine the following: |
| i. Does PPE need to be conserved? |
| ii. Does the patient have or is anticipated to have multiple IV medications that may require frequent changes? |
| iii. Does the patient require care from more than 1 nurse? |
| iv. Is there a sufficient amount of IV extension sets? |
| B. May move forward with decision-making process for positioning infusion pump outside patient room. |
| **2. Selecting and preparing correct tubing** |
| A. Use microbore tubing if possible (although large bore can be used). |
| • Using 20 feet of microbore tubing: accuracy was within specifications when infusion rates were between 5 mL and 300 mL according to ECRI.² |
| B. Use triple port connector (if needed). |
| C. Include connector volume for priming volume of extension sets. |
| D. Label tubing inside/outside room. |
| E. Use disinfection caps on all ports (do NOT use any y-sites or needle-free access ports that have been on floor). |
| F. Cover tubing on floor with orange cord protector and post signage. |
| **3. Preparing and using infusion pump** |
| A. Follow manufacturer recommendations to adjust occlusion pressure rates based on resistance, low infusion rates, and high infusion rates through CVC. |
| B. Ensure both patient and patient’s pump have wristband or patient identification taped on infusion pump. |
| C. Utilize designated infusion pump that is conducive for extension tubing; do not use syringe type pump set. |
| **4. Preparing for medications to be infused using extension sets** |
| A. Determine if medication can be infused using extension set (those that cannot include blood, fat emollients, and insulin). |
| • Consider propofol wastage if infusion outside the room. |
| B. Prime long extension sets with medication before connecting to patient to minimize risk of delayed start. |
| C. Flush rate should be the same as medication delivery rate, so medication infuses over ordered duration. |
| D. Consider compatibilities (eg, ketamine is y-site compatible with midazolam). |
| **5. Other/further actions** |
| A. Document in patient’s chart that IV pump is outside patient’s room. |
| B. Change IV administration sets and intermittent tubing every 7 days. |
| C. Develop disinfection/cleaning schedule for floor around tubing/protector. |

Abbreviations: CVC, central venous catheter; ICU, intensive care unit; IV, intravenous; PPE, personal protective equipment.

Data from ECRI.²
### Example of Step 4, Part 1 for Healthcare Failure Mode and Effects Analysis

#### TABLE 3

| Step 4, part 1: identify potential failure modes—based on the detailed outline of the process for situating IV infusion pumps outside COVID-19 ICU patient rooms |
|---|
| **1. Failure modes related to decision-making about the infusion pump outside patient room** |
| A. Nurse is unsure of current PPE and extension tubing inventory status |
| B. ICU team did not consider all of the following: PPE, patient, and tubing. |
| **2. Failure modes in selecting and preparing correct tubing** |
| A. Nurse used infusion rate of <5 mL/h (which can result in slow infusion and delayed occlusion alarm) or >300 mL/h (which can result in higher back pressure and further resulting in more frequent alarms). |
| B. Nurse does not know where to find triple port connector. |
| C. Tubing is not labeled inside and outside room. |
| D. Disinfection caps are not used on all IV tubing ports. |
| E. Tubing is not secured with orange cord protector. |
| **3. Failure modes in preparing and using the infusion pump** |
| A. Staff are unfamiliar with infusion pump manufacturer recommendations related to alarms. |
| B. Wristbands not attached to patient and/or not taped to pump. |
| C. Staff are not familiar with different pump types. |
| **4. Failure modes in preparing medications to be infused using extra extension sets** |
| A. Staff infuses blood, fat emollients, or insulin using extension tubing. |
| B. ICU team does not consider propofol wastage before infusing using an extra extension set. |
| C. Long extension set is not primed before connecting to patient. |
| C. Different flush rate than ordered is used. |
| D. Drug compatibility not confirmed. |
| **5. Failure modes related to other actions** |
| A. Placement of IV infusion pump outside patient’s room is not documented in patient’s chart. |
| B. IV administration set not changed per guideline. |

**Abbreviations:** ICU, intensive care unit; IV: intravenous; PPE, personal protective equipment.
than 20 patients in COVID-19 ICU patient rooms have had infusion pumps situated outside their rooms. Although this is a low frequency compared with some hospitals, patients were often hospitalized in the COVID-19 ICU for up to 4 weeks.

Nurses reported that it was easier to implement this process before the patient was admitted to the room. There were mixed perceptions by nurses on whether this procedure decreased PPE usage or the number of times nurses went into the room. One nurse noted that there was no delay in titrating medication infusion rates because the nurses did not have to wait until there were several patient cares or actions needed before going into the room. This was attributed to grouping care in order to reduce PPE usage. Additionally, nurses were not in 1 room for hours at a time, reducing exposure and physical strain. Nurses’ concerns included drug waste due to the extension tubing and securement of connection sites and tubing. No reports were received about infusion pump function issues if the rate was $<5$ mL/h or $>300$ mL/h.

Several failure mode themes were identified in the authors’ HFMEA. These included needing an interdisciplin ary decision, implementation of protective or preventive actions, increased risk of infection, and increased risk of human error attributed to multiple steps. The final procedure addressed these themes. The issue of drug wastage versus scarcity was important. Propofol is a high-alert medication, and tubing needs to be changed every 12 hours. This results in 34.4 mL of additional medication waste every 24 hours, based on the standard setup. For certain patients, this may be appropriate to balance PPE conservation and worker exhaustion. However, because of the national propofol shortage at the time, propofol was administered inside the ICU patient room.

There were no reported clinician or patient falls related to IV extension tubing, no identified central line-associated bloodstream infections (CLABSI), and no reported medication events related to the use of IV extension tubing. There was 1 occurrence of a propofol infusion outside the COVID-19 ICU patient room; no adverse events occurred. Last, no policies were changed for this specific procedure; it was for 1 patient care unit only and was kept with other COVID-19 special practices.

**DISCUSSION**

During a health crisis or national emergency, electronic infusion pumps may be safely situated outside ICU patient rooms to conserve resources, such as PPE or other supplies. Before implementing this procedure, stakeholders conducted an internal health and safety risk assessment, developed an approved internal procedure, completed an HFMEA, and staff instituted the practice without any adverse events reported.

The COVID-19 pandemic forced health care organizations to balance PPE preservation, health care worker exposure and absence, drug shortages, and optimal patient care simultaneously. These stressors forced the creation of strategies to conserve PPE and work with limited numbers of clinicians in ICUs and other units. Several organizations cautioned against the placement of IV pumps outside patient rooms, stating that this should be a last resort when there are critical shortages of PPE.2-6
Recent Reports About Infusion Pumps Outside COVID-19 ICU Patient Rooms

As of June 14, 2021, 5 reports related to this topic were found. The Society of Critical Care Medicine published a summary of experiences about managing patients with COVID-19 in ICUs. Situating infusion pumps outside ICU rooms was addressed extensively with recommendations for nurses and pharmacists. Almost all of the recommendations were listed in the authors’ COVID-19 ICU procedure, including securing tubing to the floor to eliminate a trip hazard. At the authors’ institution, 2 of the operating rooms nurses deployed to the COVID-19 ICU recommended use of a cord protector floor mat routinely utilized in the operating room, which was put into place.

In August 2020, an interprofessional team of ICU clinicians in the United States published their process of situating infusion pumps outside COVID-19 ICU patient rooms. Their detailed explanations of equipment and procedures provide guidance for implementing this practice. There were several differences between the authors’ procedure and what was done by Shah et al. For example, they put an identification wristband on the patient and taped a wristband to the inside of the glass door for scanning, whereas the authors’ procedure directed the nurse to put a wristband on the patient and a wristband on the infusion pump pole outside the room. With no reported adverse events using either identification method, future discussion of a preferred method with rationale may be helpful. The procedure by Shah et al was implemented institution-wide among various ICUs, although no specific outcomes were reported. With a current pandemic and limited resources, “This practice should be given consideration,” although cautions were also offered about this practice: it is not ideal and is not recommended under typical circumstances. Shah et al then studied the impact of this practice on 18 patients with COVID-19 before and after pump location inside and outside COVID-19 ICU patient rooms. They found significantly fewer nurse entries into the COVID-19 ICU rooms after pumps were situated outside the rooms (P < .0001); this was cautiously extrapolated into a corresponding decrease in PPE use. Also, there were no reports of CLABSI or extravasation for these patients, mirroring the HFMEA pump team’s experience of no reported CLABSIs.

An interprofessional team from Italy shared their experiences in situating a variety of devices outside ICU rooms during the COVID-19 pandemic. They addressed cross-contamination, technology, and regulatory issues. Information was provided for remote monitoring of a ventilator, continuous renal replacement therapy, vital signs, and drug and fluid administration. They noted that there is resistance to utilizing these remote practices in the clinical setting, but it is technologically possible. For example, when administering drugs or fluids via an electronic infusion pump, the ability to monitor pressures within the pump exists. Cautions about pump infusions were given, such as being aware of higher flow resistance attributed to longer IV tubing, knowing the internal diameter of IV tubing, and considering fluid viscosity, as well as tubing disconnection. The authors noted that it is essential to prevent transmission of infections, to keep patients safe, and to protect clinicians during a pandemic.

Last, implications for nurses, pharmacists, and providers when IV pumps are situated outside COVID-19 ICU patient rooms were summarized by an interprofessional team. They emphasized the need for more information and education when working with remotely situated IV pumps, although during a crisis time is at a premium.

Recent Reports About Related HFMEA Projects

HFMEA is a productive methodology to prevent harm to patients. However, not all reviewed HFMEA reports included outcomes. In a 2021 systematic review of 33 articles about medication safety, 31 had corrective actions recommended. Anjalee et al found that 6 articles reported a reduction in errors. In addition, the perceptions of FMEA participants about its advantages were reported; these included the ability to gather the collective knowledge of the team, analyze complex processes, increase clinicians’ awareness of health care risks, and promote safety. Several drawbacks were listed as well, such as being a time-consuming process, subjective in nature, and having a single-institution focus.

In a 2020 systematic review of 158 articles about HFMEA health care projects, Liu et al made recommendations for conducting future HFMEA projects and identified gaps in research. One of their recommendations was to have more than the usual 5 team members as reported in the literature because the complexity of health care systems often requires more experts on the team. The HFMEA pump team wanted the representation of all stakeholders involved in situating infusion pumps outside a patient’s room. This provided diverse perspectives and the ability to consider more risks. Thus, the HFMEA pump team included 10 members from 7 departments.

CONCLUSION

Situating electronic infusion pumps outside ICU patient rooms can be safe and successful but should be a practice reserved for emergencies. Proactively using a systematic method and a step by step analysis, such as an HFMEA, to analyze a new process and prevent adverse events was a productive use of time, talent, and energy in instituting a novel procedure. In addition, monitoring outcomes of high-risk process change is important to correct failures and maintain a safe system for patients. Despite many cautions, clinicians and administrators need to be prepared for future health emergencies and possible shortages. The implementation of infusion pumps outside COVID-19 ICU rooms allowed for PPE conservation, and the protocol will be in place at the authors’ institution for future pandemics, if needed. As we navigate the current pandemic and prepare...
for the future, a focus on system safety permits clinicians to focus on their patients.

Infusion therapy clinicians and managers need to be aware of issues addressed in this article, because they may be called on as stakeholders and experts for patient safety issues. Their familiarity with different patient populations, venues of care, and professional standards guiding practice make infusion therapy clinicians and managers essential to committees and councils that regularly develop and revise policies and procedures, let alone when a taskforce is needed during emergencies. Even when clinicians are not familiar with a venue of care, such as an ICU, principles of patient safety, infection prevention, use of best evidence, and methods for risk evaluation can be maintained with the necessary experts involved.

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