Sedation protocols in the pediatric intensive care unit: fact or fiction?

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Abstract: Comfort of the critically unwell pediatric patient is paramount to ensuring good outcomes. Analgesia-based, multimodal sedative approaches are the foundation for comfort, whereby pain is addressed first and then sedation titrated to a predefined target based on the goals of care. Given the heterogeneity of patients within the pediatric critical care population, the approach must be individualized based on the age and developmental stage of the child, physiologic status, and degree of invasive treatment required. In both the adult and pediatric intensive care unit (PICU), sedation titration is practiced as standard of care to meet therapeutic goals with a focus on facilitating early rehabilitation and extubation while avoiding under- and over-sedation. Sedation protocols have been developed as methods to reduce variability and optimize goal-directed therapy. Components of a sedation protocol include routine analgesia and sedation scoring with validated tools at specified intervals and a predefined algorithm that allows the titration of analgesia and sedation based on those assessments. Sedation protocols are designed to improve communication and documentation of sedation goals while also empowering the bedside team to respond rapidly to changes in a patient's clinical status. Previously it was thought that sedation protocols would consistently reduce duration of mechanical ventilation (MV) and length of stay (LOS) for patients in the PICU, however, this has not been the case. Nonetheless, introduction of sedation protocols has provided several benefits, including: (I) reduction in benzodiazepine usage; (II) improvements in interprofessional communication surrounding sedation goals and management of sedation goals; and (III) reductions in iatrogenic withdrawal symptoms. Successful implementation of sedation protocols requires passionate clinical champions and a robust implementation, education, and sustainability plan. Emerging evidence suggests that sedation protocols as part of a bundle of quality improvement initiatives will form the basis of future studies to improve short- and long-term outcomes after PICU discharge. In this review, we aim to define sedation protocols in the context of pediatric critical care and highlight important considerations for clinical practice and research.

Keywords: Clinical protocol; pediatric intensive care unit (PICU); sedatives

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Introduction

Optimizing the comfort of pediatric patients during a critical illness is an essential facet of day-to-day care in the pediatric intensive care unit (PICU). Children requiring PICU care are often subject to painful, invasive interventions that are life-sustaining. Thus, maintaining physical and psychological comfort for not only the child, but also their family and the clinical care team, is of paramount importance. Apart from non-pharmacologic options, the delivery of sedation and analgesia in this environment is considered to be the standard of care (1).

Optimum delivery of comfort in the complex ecosystem of the PICU remains elusive. A great deal has been published on (I) standardizing measures of comfort; (II) the best nonpharmacologic and pharmacologic therapies to use; (III) the best way to deliver these therapies; and (IV) how to ensure quick and appropriate responses to changes in a patient’s comfort level. However, immense variability exists and key questions remain unanswered. Should the delivery of sedation and analgesia be at an individual clinician’s discretion, or should a predefined protocol be followed that reduces subjectivity? Should determining comfort levels in the intensive care unit (ICU) be physician-, nurse-, or parent-led?

The use of defined sedation protocols has been proposed as one way to reduce variability in analgesia and sedation for critically ill patients. For the purposes of this review, we use the term sedation protocol to delineate a written, approved algorithm that is consistently utilized for patients who require continuous sedation and/or analgesia. Ideally, a standardized process could improve patient comfort, reduce subjective assessments, and improve outcomes during admission and post-ICU discharge. Despite this reasonable aim, significant controversy remains around the efficacy of such protocols. To understand why, we need to closely examine the challenges of measuring comfort, analgesia, and sedation delivery and what outcome measures are important to determine efficacy. The aim of this review will be to evaluate the potential benefit of sedation protocols and to identify targets for intervention and research.

Goals and importance of analgesia and sedation in the PICU

The aim of analgesia and sedation in the PICU is twofold: (I) to treat pain and (II) to ensure patient safety and comfort during invasive treatments (1). Other possible goals include optimizing patient-ventilator synchrony, reducing oxygen demand, line and tube maintenance, and reducing anxiety (2,3). Although every patient requires adequate analgesia for noxious stimuli (e.g., the endotracheal tube), sedative needs may vary widely from patient to patient depending on the goals of care.

As such, the pendulum for ideal depth of sedation and analgesia has swung back and forth over time. Initially, it was widely held that young children and infants did not experience the same degree of pain or discomfort as adults. As a result, pediatric patients often received minimal sedation and analgesia. Poorer outcomes were then noted with undertreated pain and anxiety, including delayed healing and increased stress on patients and caregivers (4). With these findings, children were subsequently more deeply sedated with a variety of agents, most commonly benzodiazepines and opioids (5). Increased sedation resulted in a new set of problems with clinical impact, including prolonged mechanical ventilation (MV) (6), increased iatrogenic withdrawal syndrome (IWS) (7,8), delirium (9), and the potential negative ramifications of these medications on immature and developing brains (8,10). Given the increasingly deleterious effects of over-sedation, the most recent shift in sedation goals has been to titrate sedation to minimum but effective targets, with an emphasis on early extubation and rehabilitation while maintaining optimal analgesia.

Consensus guidelines from the United Kingdom (3) and the European Society of Pediatric and Neonatal Intensive Care (ESPNIC) (11) have all advocated similar recommendations. Firstly, pain should be assessed with age-appropriate validated pain scales to provide consistency between clinicians and allow for evaluation of an intervention’s effect. Commonly used pain assessments in the PICU (10) include: (I) COMFORT Score, (II) Faces, Legs, Activity, Cry and Consolability (FLACC) score, and (III) Multidimensional Assessment of Pain Scale. Pain assessments should be performed regularly, particularly if the child is on an analgesic infusion, and in conjunction with patient and family assessment (11). Thus, a key component of sedation management in the PICU is that pain should be treated before targeting sedation depth, i.e., analgesia-based sedation. Clinical cues for pain or agitation from children have significant overlap, but attempting to distinguish between them allows clearer setting of goals and choice of therapies.

Secondly, an optimal sedation goal should be determined for the patient by carefully considering clinical status and
disease severity and using validated sedation scoring tools (3,11). Achieving a targeted sedation level can often be challenging as it depends on the child’s neurodevelopmental stage, illness severity, and by association the degree of invasive support that is required. For example, some patients may require complete immobility with deep sedation for physiologic stability, whereas others may tolerate MV and endotracheal intubation with only opioid therapy. Once the desired level of sedation has been established, it should be assessed regularly with validated scales (3). Sedative doses should then be titrated based on the assessed depth of sedation and adjusted accordingly.

**Sedation assessment tools**

A central component of all sedation protocols is the ability to measure a patient’s sedation level at regular intervals using a reproducible, systematic, and validated approach. Utilization of standardized assessment tools enables consistent communication between PICU team members and prevention of over- or under-sedation (1,11).

However, there is no gold standard tool for achieving target sedation goals (12). The challenge of such a tool in pediatrics is that it must be validated for a wide range of ages and developmental stages. Additionally, such tools need to fit within local work practices and be reproducible between various clinicians. The interval for regular assessments is defined by the scale being used and the protocol in place, but general recommendations are that it be completed every 4 to 8 hours or as indicated by the child’s clinical condition (11).

Currently, the three validated pediatric sedation assessment tools for children supported on MV are: (I) the State Behavioral Assessment Scale (SBS); (II) the COMFORT Scale and its modification, the COMFORT-B Scale; and (III) the Richmond Agitation Sedation Scale (RASS) (Table 1). The SBS provides a numeric rating for a standardized range of defined behaviors in response to voice, gentle touch, or noxious stimuli (13). The scale ranges from agitated (+2) to unresponsive (−3) (13). The COMFORT Scale (14) was created to evaluate distress in eight domains: alertness, calmness/agitation, respiratory...
response, physical movement, mean arterial blood pressure (MAP), heart rate, muscle tone, and facial tension. These domains are evaluated on a Likert scale from 1 to 5, from which patients receive a cumulative score (14). Given concerns that physiologic variables are influenced by medications administered in the PICU, the COMFORT-B Scale was developed and validated to eliminate physiologic variables from the score (15). The RASS is a responsiveness scale that assesses sedation and agitation and was recently validated for use in critically ill children. It is a single value numeric score that ranges from −5 to 4 to describe a patient’s behavior from unarousable to combative in response to voice and touch. The RASS is unique in that it is the only sedation score that incorporates eye contact in response to verbal stimuli (16). No studies have directly compared these three validated sedation assessment tools.

**Components and goals of a sedation protocol**

Sedation protocols are used to achieve goal-directed, patient-specific sedation and analgesia for each clinical scenario while minimizing both under- and over-sedation. Frequently, physicians use nonstandardized practices to titrate sedation in critically ill children, leading to significant practice variation in terms of drug choice, method of drug delivery, and adherence to protocols (5). Clinicians and researchers anticipated that a formulated algorithm incorporating validated scores for sedation and pain and the use of regular assessments to guide medication titration would provide substantial benefit. Two types of standardized sedation strategies have been studied in critically ill children: protocol-directed sedation and daily sedation interruption. Protocol-directed sedation is ordered by a physician with specific goals for a patient’s sedation level and is implemented by a member of the healthcare team, often bedside nursing. The patient’s sedation level is assessed on a regular time interval with a validated sedation assessment tool, and sedative agents are titrated to achieve the targeted sedation goal (17). Such protocols allow the interprofessional team to set goals and transfer decision-making authority for analgesia and sedation to bedside clinicians, usually nurses (18). Daily sedation interruption requires a period of time each day that a patient’s sedation is paused or decreased to achieve a goal level of alertness defined by a sedation assessment tool (19). The aims of sedation interruption are to reduce the total amount of sedation exposure and facilitate early weaning from the ventilator.

The reported benefits of a sedation protocol are wide-ranging. Sedation protocols introduce standardization of assessments of pain and sedation (10), empower bedside clinicians to respond to those assessments, and allow for a common language to be spoken regarding analgesia and sedation within the interdisciplinary team (18,20). These protocols give priority to the issue of a child’s comfort. Other reported benefits are more ambitious, including improvements in duration of MV and reduced ICU length of stay (LOS) (21). Although many benefits of sedation protocols have been reported, they have not always been supported by strong evidence. The reported benefits are mostly extrapolated from adult studies and are listed in Box 1.

**Box 1 Reported benefits from implementation of a sedation protocol**

- Decreased duration of MV
- Decreased ICU LOS
- Decreased hospital LOS
- Decreased incidence and severity of delirium
- Earlier mobilization
- Reduced over- and under-sedation
- Decreased sedative and opioid doses
- Decreased incidence of iatrogenic drug withdrawal
- Improved communication of sedation goals in the interdisciplinary team
- Improved documentation of sedation goals

MV, mechanical ventilation; ICU, intensive care unit; LOS, length of stay.

**What is the evidence for a sedation protocol?**

In an international survey of PICUs, sedation scoring tools were used in 70% of PICUs, but only 42% reported using them as part of routine daily rounds and patient care goals (5). Only 27% of PICUs had written sedation protocols with treatment algorithms in place (5). In the adult ICU literature, the results of protocol-directed sedation studies are conflicting. Individual adult studies demonstrate reductions in duration of MV, ICU LOS, and mortality (22-24); however, a large systematic review and meta-analysis of protocol-directed sedation did not
reproduce the same benefit (17).

The largest study in the pediatric population to assess protocol-directed sedation is the Randomized Evaluation of Sedation Titration for Respiratory Failure (RESTORE) trial (25). RESTORE was an unblinded, multicenter, cluster-randomized, prospective clinical trial conducted between 2009 and 2013 that enrolled 2,449 pediatric patients receiving MV from 31 US PICUs. PICUs were randomized into either the study group or a control group. The study group included a nurse-titrated sedation protocol that incorporated prescriber-directed SBS, regular arousal assessments utilizing SBS, and daily extubation readiness tests. PICUs in the control group continued to manage sedation in their usual nonstandardized way. The primary outcome, duration of MV, did not differ between the groups. There were also no differences in PICU LOS, hospital LOS, in-hospital mortality, inadequate sedation management, or clinically significant IWS. In a secondary exploratory analysis, patients in the intervention group were exposed to fewer classes of sedative agents (2 vs. 3 agents) and fewer cumulative days of opioid administration (9 vs. 10 days). The percentage of study days in which patients were awake and calm while intubated was higher in the intervention group (86% vs. 75%), though the intervention group had a greater percentage of days with a higher pain score (50% vs. 23%) and higher SBS (60% vs. 40%). Additional findings included an increase in post-extubation stridor and immobility-related pressure ulcers in the control group of patients. Reducing these potential sedation-related events could be considered a potential benefit of protocolized sedation. Ultimately, the authors concluded that a nurse-implemented, goal-directed sedation protocol did not reduce the duration of MV from that with usual care (25), but that children can be more awake and calm without harm. In a subsequent follow-up study of 1,360 patients, the authors evaluated post-discharge outcomes of children who were randomized to a sedation protocol or usual care. There were no difference in functional status, quality of life, or indication of post-traumatic stress disorder between the groups. The authors concluded that a sedation strategy in which patients are more awake does not produce long-term harm (26).

Much of the remaining evidence regarding sedation protocols in pediatrics has come from before-and-after protocol implementation studies with different outcome measures (18,20,21,27-33) (Table 2). Similar to the adult literature, these studies have produced mixed results with no clear benefit. Prior to the RESTORE study, Poh and co-authors (34) completed a systematic review of sedation protocols in pediatric critical care. The driver for the review was that many societies had included sedation protocols in their standard recommendations to improve sedation practice and standardize care with poor supportive evidence. Within the studies reviewed by Poh et al. (34), some reported benefits included decreasing unplanned extubation, IWS, and sedation doses and duration. There were no consistent positive results in terms of mortality, duration of MV, or LOS. Most studies were single center, and hence applicability to other PICUs was difficult. Another challenge was the lack of a clear definition for sedation-related adverse events.

No evidence in the pediatric literature has shown that the use of a protocol reduces duration of MV or PICU LOS. Most studies, including the large RESTORE study (25), have shown no difference in these outcome measures (25,27-29). The exceptions were one Korean study, which showed a decrease in duration of MV (21), and an Australian study, which demonstrated an increase in duration of MV (31). In the latter study, the authors reported a significant number of younger children in the post-implementation phase who were less likely to be extubated early post-cardiac surgery. That group potentially influenced their result. Aitken et al. (17) examined randomized controlled trials (RCT) of protocol-directed sedation in both adults and children, including three adult studies and the pediatric RESTORE study. All of these studies compared protocol-directed sedation with usual care and found no evidence of difference in MV duration and no clear evidence of difference in ICU mortality, hospital mortality, ICU LOS, or unplanned extubations.

**Are we asking the right questions about sedation protocols?**

The pediatric literature on sedation protocols leads us to ask: are MV duration and PICU LOS the right outcomes to assess when examining the benefit of minimal but effective sedation? Both of these variables are discrete time measurements that are affected by many factors not captured in the studies and that may be difficult to measure. For example, ventilation practice, hospital throughput, staffing ratios, and clinician preferences can all influence these outcome measures and hence make it difficult to show a benefit related to changes in sedation. In addition, all of the studies compared the use of a sedation protocol to
Table 2: Review of before-and-after sedation protocol implementation studies and reported outcomes

| Author          | Sedation assessment | Duration of MV | PICU LOS | Sedation-related adverse events† | IWS               | Sedation doses | Sedation assessments |
|-----------------|---------------------|----------------|----------|----------------------------------|-------------------|----------------|----------------------|
| Dreyfus et al.  | COMFORT-B           | Unchanged      | Unchanged| No                               | Trend towards decrease | No difference  | Increase in target sedation scores. Decrease in over-sedated scores |
| (27)            |                     |                |          |                                  |                   |                |                      |
| Gaillard-Le Roux et al. (28) | COMFORT-B           | Unchanged      | Unchanged| No                               | No difference     | Decrease in midazolam and ketamine dose | No difference |
| Neunhoeffer et al. (29) | COMFORT-B           | Unchanged      | Unchanged| No                               | Decreased         | Decrease in benzodiazepine dose | Increase in sedation assessments. More episodes of under-sedation reported |
| Ist et al. (30) | COMFORT-B           | N/A            | N/A      | N/A                              | Increase in morphine and midazolam dose | Increase in target sedation scores |                      |
| Larson and McKeever (31) | COMFORT-B           | Increased      | N/A      | N/A                              | Decrease in midazolam dose; decrease in morphine infusions, with increase in bolus doses | Increase in frequency of sedation assessments |                      |
| Alexander et al. (18) | Behavioral cues    | N/A            | N/A      | N/A                              | Increased sedation received on protocol | Decreased under-sedation |                      |
| Larson et al. (20) | COMFORT-B           | N/A            | N/A      | N/A                              | Decrease in ketamine | Increase in sedation assessments and in documentation and communication of sedation goals |                      |
| Deeter et al. (33) | Seattle PICU Comfort Score | Unchanged      | Unchanged| Decrease in accidental extubation | –                  | Reduced days of sedatives | Not reported |
| Jin et al. (21) | COMFORT             | Decreased      | Decreased| N/A                              | Decreased         | Decrease in sedative doses | Not reported |

†, sedation-related adverse events include unplanned extubation, line removal, and ventilator-associated pneumonia. MV, mechanical ventilation; PICU, pediatric intensive care unit; LOS, length of stay; IWS, iatrogenic withdrawal syndrome; N/A, not assessed.
“usual” or “standard” care. However, standard care has no consistent definition. An important consideration regarding standard care is that mortality in pediatric critical care is very low (35) and outcomes are improving. It is possible that “standard care” in pediatric critical care has advanced to a point that a meaningful benefit cannot be shown in the outcomes measured.

Nonetheless, some benefits of sedation protocols have been shown. One positive outcome that was demonstrated in several studies was the reduction in use of benzodiazepines, midazolam in particular, with no increase in harm to patients (28,29,31,33). It is possible that the implemented protocols directed clinicians away from midazolam use and hence is an expected result. The studies also showed that using less midazolam in the PICU was not harmful and resulted in higher success in meeting sedation targets (31) without an increase in adverse effects. Literature to support minimization of benzodiazepines is useful, as growing evidence suggests that benzodiazepines can have a negative impact on cognitive development and incidence of delirium (8,10).

Studies have shown mixed results with regard to overall analgesic and sedative exposure. Several studies have shown an increase in sedative exposure with the implementation of protocolized sedation (18,30,32), thought to be secondary to improvement in target sedative goals and reduction in under-sedation. Conversely, other studies have shown a reduction in total sedative exposure (27–29), therefore reducing over-sedation. Without knowledge of an institution's sedation practice prior to the implementation of a protocol, it is difficult to interpret the impact on cumulative sedative or analgesic dosing, as it very much reflects cultural practices within the unit; however, achieving better target sedation goals is promising.

Use of sedation protocols improves compliance with sedation assessments (27,30,31) and achievement of target sedation scores within the prescribed range. An improvement in communication and documentation of sedation assessments has also been reported (20). In a survey of nurses and junior and senior physicians, all respondents reported that the introduction of a sedation protocol facilitated team decision making and resulted in more efficient and rapid intervention to ensure patient comfort (18). Another important benefit of sedation protocols is implied rather than actually measured in the studies reported. That is, the benefit of empowering the bedside clinician, generally the nurse, to act on the assessment of their patient’s comfort cannot be understated.

It is generally accepted that a routine part of nursing care is to complete assessments of each patient's comfort level. Without the predefined protocol, nurses would need to first find a physician, who would then assess the patient and prescribe dose adjustments to address comfort, causing a real time delay in delivering comfort to the patient. With a sedation protocol, the bedside clinician can respond immediately to address the patient's comfort by working within predefined boundaries. This approach allows for the timely communication, documentation, and achievement of target sedation goals. This advantage is not explicitly reported in the studies but rather implied. Consideration should be given to capturing this benefit in future study designs.

What about daily sedation interruption?

Another sedation protocol that gained popularity initially in adult critical care is the concept of daily sedation interruption to minimize cumulative sedation and thereby decrease MV duration and ICU LOS (36). The results of these studies in the adult literature are mixed. Several studies showed decreases in duration of MV and total cumulative dose of sedatives administered (37,38); however, a larger more recent RCT showed no decreases in duration of MV or ICU LOS when protocolized sedation was combined with daily sedation interruption (39).

To date, three pediatric studies have investigated daily sedation interruption. The first two were single-center RCTs that compared daily sedation interruption to usual care. Both studies demonstrated decreased length of MV, decreased use of sedation, and shorter PICU LOS (40,41). The largest study was a multicenter RCT that compared daily sedation interruption and protocol-directed sedation. The study included three tertiary hospitals in the Netherlands and 129 patients. The authors found no differences between the two groups in length of MV, cumulative dose of benzodiazepines, or PICU LOS, and daily sedation interruption was associated with increased mortality and need for reintubation (42). Thus, unlike protocolized sedation, daily sedation interruption has not been widely adopted in the pediatric population because of the potential for adverse events.

Role of sedation protocols in preventing IWS in the PICU

It is well documented that prolonged use of analgesics and
sedatives can lead to tolerance and dependence and hence IWS if those medications are discontinued abruptly (11). Establishing a sedation protocol could potentially prevent IWS by reducing the total cumulative doses of medications or by building an active weaning process into the protocol as guided by sedation targets. Validated withdrawal scores are available for use in pediatrics. These include the Withdrawal Assessment Tool-1 (WAT-1) and the Sophia Observational Withdrawal Symptoms scale (11). In one study, introduction of a sedation protocol significantly reduced the incidence of IWS from 23.6% to 12.8% (29). Other studies have reported trends toward reduction of withdrawal symptoms after introduction of a sedation protocol (21,27). In the RESTORE study, no difference was seen in the incidence of IWS (25). Despite mixed results, prevention of IWS is an important consideration for future studies.

### Role of sedation protocols within PICU liberation bundles

As mortality in the PICU has decreased, several quality improvement initiatives have been aimed at mitigating the long-term psychologic, social, and physical impairment of critical illness (43). ICU liberation bundles, also known as ABCDEF Bundles (44) (Table 3), are evidence-based guidelines to liberate patients from the harmful effects of a PICU stay (10,43). It should be noted that sedation protocols are a critical element of these bundles, directly addressing “A” and “C” and having an impact on “B”, “D”, and “E”. Special consideration should be given to the screening and diagnosis of delirium (“D” in the liberation bundle) when utilizing sedation protocols. A number of delirium symptoms overlap with those observed with pain, distress and withdrawal, thus integration of validated delirium screening is essential. There are two validated commonly used bedside screening tools for delirium in pediatric intensive care patients: The Pediatric Confusion Assessment Method for the ICU (pCAM-ICU or psCAM-ICU for preschool-age children) and the Cornell Assessment of Pediatric Delirium (CAPD) (10,11). A review of the diagnosis and management of delirium is outside the scope of this review, but future research must assess sedation protocols as a component of multifaceted strategies to improve PICU outcomes.

### Barriers and sustainability of sedation protocols

Institutional implementation of sedation protocols through quality improvement initiatives has been associated with numerous challenges. In one study, Yaghmai et al. (32) aimed to evaluate whether the initial benefits of their institution’s sedation protocol persisted after a 4-year period. When the sedation protocol tool was initially introduced in 2008, implementation reduced LOS, sedation days, and number of opioid infusion days and decreased benzodiazepine use (33). However, the only benefit maintained was the reduction in benzodiazepine dose (32). The authors concluded that sustainability is an important consideration when creating a sedation protocol and that it requires clinical champions to carry it forward, a robust education program, ongoing quality assessment, and consistent audits (32). Other barriers to implementation have also been described, including lack of agreement and familiarity with protocol design (18,30) and challenges with unit culture change around sedation minimization (30). Reluctance by staff to reduce sedation at night caused one institution to halt protocol use overnight (30). Barriers to implementation and sustainability of sedation protocols are difficult to study. Additionally, it is challenging to capture

| Bundle component | Description |
|------------------|-------------|
| A                | Assess, prevent, and manage pain |
| B                | Spontaneous awakening trials and spontaneous breathing trials |
| C                | Choice of analgesia and sedation |
| D                | Delirium: assess, prevent, and manage |
| E                | Early mobility and exercise |
| F                | Family engagement and empowerment |

PICU, pediatric intensive care unit.

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Table 3 PICU liberation bundle (44)
reasons behind deviations from a protocol in the traditional study designs used to date. Adherence to protocol and preservation of protocol integrity over time are important considerations for future studies.

**Challenges with research in sedation**

As we have described above, there are many challenges associated with conducting research to better understand sedation in critical care (36). A high degree of individual variability is exacerbated by the wide age range and differences in pharmacokinetics and pharmacodynamics in the pediatric population. The variability in intensity of nursing and medical bedside care and drugs that are available internationally also adds to the challenge of studying this topic (36). The success of a sedation minimization strategy depends on factors influenced by unit culture, including local practices, nurse-to-patient ratios, and intensity of training and experience (36), which is not always translatable or easy to study. Designing studies to address these factors can also be challenging because of the multicomponent structure of such an intervention (45), including patient monitoring and assessment, measurement and evaluation of sedation, practitioner education and training, and interprofessional collaboration (45). Finally, the most commonly used study designs have all compared sedation protocols to “usual care”, a term that implies a uniform practice standard (45). In reality, though, the substantial variability and inconsistency in usual care makes any comparison challenging to study.

**Conclusions**

Strong evidence from available literature supports the premise that pediatric patients can safely tolerate minimal effective sedation with the use of a sedation protocol. No increases in adverse outcomes have been attributed to a decrease in sedation. Other reported benefits include improved communication, targeting of sedation goals, reduction of benzodiazepine use, and a reduction in IWS. A review of the evidence, however, does not reveal a clear benefit to protocol-directed sedation or daily sedation interruption over usual care in duration of MV, LOS, or mortality. Nevertheless, protocolized sedation is gradually becoming part of the accepted milieu of the PICU, as it can facilitate interventions that do impact PICU outcomes such as early mobility, reduction in delirium, and family interaction. Future studies should consider the incorporation of targeted sedation delivery in the context of bundled care in the assessment of PICU outcomes.

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