Sedation protocols to reduce duration of mechanical ventilation in the ICU: a Cochrane Systematic Review

Leanne M. Aitken, Tracey Bucknall, Bridie Kent, Marion Mitchell, Elizabeth Burmeister & Samantha Keogh

Aim. Assess the effects of protocol-directed sedation management on the duration of mechanical ventilation and other relevant patient outcomes in mechanically ventilated intensive care unit patients.

Background. Sedation is a core component of critical care. Sub-optimal sedation management incorporates both under- and over-sedation and has been linked to poorer patient outcomes.

Design. Cochrane systematic review of randomized controlled trials.

Data sources. Cochrane Central Register of Controlled trials, MEDLINE, EMBASE, CINAHL, Database of Abstracts of Reviews of Effects, LILACS, Current Controlled Trials and US National Institutes of Health Clinical Research Studies (1990–November 2013) and reference lists of articles were used.

Review methods. Randomized controlled trials conducted in intensive care units comparing management with and without protocol-directed sedation were included. Two authors screened titles, abstracts and full-text reports. Potential risk of bias was assessed. Clinical, methodological and statistical heterogeneity were examined and the random-effects model used for meta-analysis where appropriate. Mean difference for duration of mechanical ventilation and risk ratio for mortality, with 95% confidence intervals, were calculated.

Results. Two eligible studies with 633 participants comparing protocol-directed sedation delivered by nurses vs. usual care were identified. There was no evidence of differences in duration of mechanical ventilation or hospital mortality. There was statistically significant heterogeneity between studies for duration of mechanical ventilation.

Conclusions. There is insufficient evidence to evaluate the effectiveness of protocol-directed sedation as results from the two randomized controlled trials were conflicting.
Introduction
Sedation management of critically ill patients is a core component of critical care; these patients are often treated with invasive and difficult-to-tolerate procedures and treatments. Ensuring comfort throughout this process assists recovery and ensures humane treatment (Mehta et al. 2009). To promote this, appropriate sedation is essential for all critically ill patients, as is associated pain relief and anxiolysis. To support this practice, a systematic review of randomized controlled trials (RCTs) that examined the effectiveness of nurse-directed sedation protocols was undertaken and published as a Cochrane systematic review (Aitken et al. 2014); this current paper presents a summarized version of that review.

Background
Growing evidence suggests that sedation is poorly managed; one systematic review of 36 studies found a substantial incidence of sub-optimal sedation (Jackson et al. 2009). The detrimental impact of poor sedation practices extends from under-sedation to over-sedation. Under-sedation has the potential to lead to agitated patients with compromised long-term psychological recovery, while over-sedation may lead to increased intensive care unit (ICU) and hospital lengths of stay and poor long-term recovery (Mehta et al. 2009). There is some evidence to suggest links between short-term measures (such as intensive care and hospital lengths of stay) (Kollef et al. 1998, Schweickert & Kress 2008, Jackson et al. 2010), adverse events (such as self extubation) (Girard et al. 2008) and long-term outcomes such as ICU memory recall and psychological recovery (Ringdal et al. 2006, Samuelson et al. 2006, Jackson et al. 2010).

Sedation refers to the administration of pharmacological agents designed primarily to induce a sedative effect in patients. Sedation does not include pharmacological agents administered primarily for other reasons, such as analgesics, although these agents might have some secondary sedative effect. Internationally, there is a range of different methods of managing patients’ sedation needs.

Various strategies have been proposed to improve sedation management of critically ill patients: sedation assessment instruments (Riker et al. 1999, Ely et al. 2003, Curley et al. 2006); sedation guidelines, algorithms or protocols to guide assessment and therapy (Jacobi et al. 2002, Sessler & Pedram 2009); implementation of daily sedation interruptions (Kress et al. 2000); targeting minimal levels of sedation and regular assessment of sedation and analgesia requirements (Schweickert & Kress 2008). Despite a core component of many of these recommendations being the use of an algorithm or protocol, there is evidence to suggest that
sedation guidelines remain poorly implemented, with less than 50% of critical care units in Canada, US and Denmark indicating such use (Sessler & Pedram 2009). This lack of implementation may be due to the inconsistent results in the studies examining the effect of protocol-directed sedation (Brook et al. 1999, De Jonghe et al. 2005, Elliott et al. 2006, Quenot et al. 2007, Bucknall et al. 2008). Protocol-directed sedation is ordered by a physician, contains guidance regarding sedation management and is implemented by nurses, pharmacists or other members of the healthcare team. Selection of the most appropriate sedative agent, and when to commence, increase, decrease or cease administration of the agent, is based on patient assessment, usually with the aid of a sedation scale. Protocols may include an analgesic component (Brook et al. 1999). Protocol-directed sedation is distinct from, but related to, protocol-directed weaning, which is specifically directed towards limiting the duration of mechanical ventilation (Blackwood et al. 2014).

Use of a protocol may improve sedation by incorporating regular patient assessment with planned changes to sedative or analgesic agents, or both. There is widespread evidence of international variation in sedation assessment and management practices (Mehta et al. 2009, O’Connor et al. 2009). The potential to reduce the individual clinician variation is statistically significant, with management based on standardized assessment practices. Despite widespread use of sedation protocols there is mixed evidence as to their effectiveness.

The review

Aims

The aim of this study was to assess the effects of protocol-directed sedation management on the duration of mechanical ventilation and other relevant patient outcomes (Table 1) in mechanically ventilated ICU patients.

Design

RCTs and quasi-randomized controlled trials published in any language were included. An RCT was defined as a study, where patients were allocated to treatment groups based on a random or quasi-random method (e.g. using random number tables, hospital number, date of birth).

All ICU patients who were mechanically ventilated (via endotracheal or tracheostomy tube) were included. If eligible studies had included both patients who met the above criteria and those who did not, data were excluded unless the subpopulations were reported, or able to be obtained.

Table 1 Primary and secondary outcomes for systematic review.

<table>
<thead>
<tr>
<th>Primary outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Duration of mechanical ventilation measured in hours for the entire duration of the first ICU stay for each patient</td>
</tr>
<tr>
<td>2. ICU and hospital mortality</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Secondary outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. ICU length of stay</td>
</tr>
<tr>
<td>2. Hospital length of stay</td>
</tr>
<tr>
<td>3. Total dose of sedation</td>
</tr>
<tr>
<td>4. Adverse events (e.g. non-planned extubation)</td>
</tr>
<tr>
<td>5. Incidence of delirium</td>
</tr>
<tr>
<td>6. Memory function</td>
</tr>
<tr>
<td>7. Psychological recovery</td>
</tr>
<tr>
<td>8. Cognitive recovery</td>
</tr>
<tr>
<td>9. Quality of life</td>
</tr>
<tr>
<td>10. Incidence of tracheostomy</td>
</tr>
</tbody>
</table>

ICU, intensive care unit.

The target intervention was protocol-directed sedation management which was compared with non-protocol-directed sedation management. Protocol-directed sedation was defined as sedation directed by a protocol or algorithm that was ordered by a medical officer, contained guidance regarding sedation management and was implemented by nurses, pharmacists or other members of the healthcare team with sedation increased or decreased based on patient assessment. The guidance regarding sedation management consisted of a series of decision points or decision algorithms that assisted clinicians to make decisions regarding increasing, decreasing or maintaining current sedation levels. Protocols included provision for administration of analgesics in addition to sedative agents. Medical officers may have continued to be involved in sedation assessment and management beyond the point of ordering the sedation protocol, but any protocol that required physician approval for changes in amounts of sedation was excluded. The essential element of protocol-directed sedation was that other members of the healthcare team could alter the level of sedation being administered without consulting with a medical officer. Usual care was defined as physician-led sedation management of mechanically ventilated patients according to local practice, where no specific strategies were implemented to change the level of sedation. Sedative agents may or may not have been different to those used in the intervention; importantly the intervention was not about the agents that were used but how they were used.

Search methods

The Cochrane Central Register of Controlled trials (CENTRAL) (2013, Issue 11), MEDLINE (OvidSP; from
1990–November 2013), EMBASE (OvidSP; from 1990–November 2013), CINAHL (BIREME host; from 1990–November 2013), Database of Abstracts of Reviews of Effects (DARE) (from 1990–November 2013), LILACS (1990–November 2013), Current Controlled Trials and US National Institutes of Health Research Studies (from 1990–November 2013) were searched. An example of the search strategy is provided in Supplementary Table 1. The search was re-run in October 2014 and any studies of interest will be dealt with when the review is updated. The MEDLINE search strategy was combined with the Cochrane highly sensitive search strategy, as detailed in Higgins and Green (2011) and was adapted for searching all other databases. Relevant critical care journals, reference lists of identified published trials, abstracts of relevant conference proceedings and the reference lists of relevant articles were hand-searched to identify further clinical trials. Relevant trial authors were contacted to identify any additional studies. We searched specific websites for relevant ongoing trials:

1. International Clinical trials registry (www.who.int/trialssearch);
2. International Standard Randomized Controlled Trials (www.controlled-trials.com/isrctn);
3. Country specific trial websites for the UK, South Africa, India, Hong Kong, China and Australia and New Zealand.

No language restriction was imposed.

Quality appraisal

Two authors (LA and TB or MM) independently assessed the methodological quality of each eligible trial as per the Cochrane Collaboration guidelines (Higgins & Green 2011); disagreements were resolved by discussion. Where potential conflicts of interest existed, the relevant author was excluded from the process. Seven domains were assessed to determine risk of bias (Table 2); we considered a trial as having a high risk of bias if one or more of the assessment domains was rated as high risk or unclear.

We assessed clinical heterogeneity for key participant and sedation protocol characteristics. Study cohorts were considered sufficiently similar for participant and intervention characteristics to suggest data could potentially be pooled for statistical analysis. We assessed statistical heterogeneity using the $I^2$ statistic. Where this analysis suggested statistical heterogeneity was moderate or greater, we have noted that care should be taken when interpreting the results for that outcome. In the absence of sufficient homogeneity between the studies, we provided a descriptive presentation of the results. We did not undertake meta-regression due to insufficient studies and appropriate homogeneity; similarly there were insufficient studies (<10) to construct a funnel plot to explore the symmetry of the intervention effects to assess for publication bias.

Data abstraction

Two authors (LA and TB) independently reviewed all titles and decided on the inclusion of studies based on selection criteria, then extracted standardized data from each study. We resolved differences and avoided conflicts by consulting a third author (MM). If a study had insufficient data to complete data extraction or if we required data clarification, we contacted the authors of the study. We considered the studies to have sufficient data if at least one of the listed outcomes (either primary or secondary) was reported.

Synthesis

Subject to the absence of clinical heterogeneity, we undertook an analysis using Review Manager 5 software (RevMan 2013). For continuous data, the mean difference (MD), or standardized mean difference (SMD) and 95% confidence interval (CI) for summary statistics (hospital and ICU length of stay, duration of mechanical ventilation) was used wherever possible. We found the data to be skewed and, due to the unavailability of source data related to one study, we were unable to transform the data for analysis. For dichotomous data, we used risk ratio (RR) and 95% CI.

We used the results of intention-to-treat (ITT) analyses for all analyses so all data extracted reflected the original allocation group. There was no evidence of multiple observations or outcome measurements in either of the included studies and all outcome measurements were taken at the same time point in both studies. The duration of mechanical ventilation was measured on the same group of patients throughout their ICU stay. Both included studies had a small number (less than 4%) of participants, who were recruited into the studies despite not meeting inclusion criteria (re-admission to ICU, patient awaiting rapid transfer to another ICU) and we excluded these patients from all analyses. Published study reports identified complete data for all included participants, indicating there were no drop-outs in either study.

If studies were sufficiently homogenous, we planned to conduct a meta-analysis using a fixed-effect model or where heterogeneity existed, a random-effects model. We conducted meta-analyses for all outcomes where possible, although the meta-analyses for many of the outcomes should be interpreted with caution due to the presence of substantial
Table 2 Characteristics, strengths and limitations of included studies.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Brook et al. (1999)</th>
<th>Bucknall et al. (2008)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intervention (see Supplementary Table 2 for more detail)</td>
<td>RCT, 322 patients in a closed medical ICU in a university-affiliated teaching hospital in US.</td>
<td>RCT, 316 patients in a closed general ICU in a metropolitan teaching hospital in Australia.</td>
</tr>
<tr>
<td>Protocol-directed sedation vs. non-protocol-directed sedation (usual care). Sedation protocol required nurses to determine type, method of administration and dosage of analgesics and sedatives after assessing using the Ramsay Scale.</td>
<td>Protocol-directed sedation vs. non-protocol-directed sedation. Sedation protocol required nurses to determine the type, method of administration, dosage of sedation or analgesia after assessing using the Sedation-Agitation Scale.</td>
<td></td>
</tr>
<tr>
<td>Outcomes measured (see Supplementary Table 2 for more detail)</td>
<td>Primary outcome – duration of mechanical ventilation. Secondary outcomes – ICU and hospital LOS, hospital mortality, rates of organ failure, re-intubation and tracheostomy.</td>
<td>Primary outcome – duration of mechanical ventilation. Secondary outcomes – ICU and hospital LOS, ICU and hospital mortality, rates of self extubation and tracheostomy.</td>
</tr>
<tr>
<td>Assessment of bias (see Supplementary Table 3 for more detail)</td>
<td>Generally low risk of bias with the exception of the following:</td>
<td>Generally low risk of bias with the exception of the following:</td>
</tr>
<tr>
<td>- Unclear risk of selection bias due to randomization process</td>
<td>- High risk of performance bias due to inability to blind participants and personnel</td>
<td>- Unclear risk of other bias due to lack of description of some aspects of usual care</td>
</tr>
</tbody>
</table>

RCT, randomized controlled trial; ICU, intensive care unit; LOS, length of stay.

heterogeneity (duration of mechanical ventilation, length of ICU stay and incidence of tracheostomy). Analyses were considered significant at the alpha = 0.05 level. Estimates of precision were assessed by interpretation of CIs, such as widths, overlapping and inclusion of the null hypothesis.

Intensive care patients were a heterogeneous group. Given the small number of studies and limited variation in the included participants and methods, we could not undertake sub-group or sensitivity analyses.

We used the principles of the GRADE system to assess the quality of the body of evidence associated with outcomes reported (Guyatt et al. 2008). The GRADE approach appraises the quality of evidence based on the extent to which one can be confident that an estimate of effect or association reflects the item being assessed. The quality of a body of evidence considers within-study risk of bias (methodological quality), the directness of the evidence, heterogeneity of the data, precision of effect estimates and risk of publication bias.

Results

The results of the search and selection of studies are summarized in the PRISMA study flow diagram (Figure 1). After exclusion of duplicates, we identified 2041 records, with 21 full-text articles retrieved. We excluded 13 of these as they did not address our research question, for example, they answered different questions or provided a review of the topic and we excluded six studies as, although they addressed the question of our review, they did not use a randomized or quasi-randomized design (Brattebo et al. 2002, De Jonghe et al. 2005, Elliott et al. 2006, Quenot et al. 2007, Arias-Rivera et al. 2008, Tobar et al. 2008). We re-ran the search in October 2014. We identified a further 482 records after removing duplicates; we identified one study of interest and we will report this study when we update the review. We included two studies (Brook et al. 1999, Bucknall et al. 2008). The studies were similar in design and examined the impact of protocol-directed sedation on a range of outcomes including duration of mechanical ventilation, mortality, ICU and hospital length of stay and some adverse events (Table 2).

Brook et al. (1999) enrolled 332 participants from a single 19-bed medical ICU in a university-affiliated urban teaching hospital in the USA, with data collected in 1997–1998. In contrast, Bucknall et al. (2008) enrolled 316 participants from a 24-bed mixed ICU in a major Australian metropolitan university-associated teaching hospital. Participants were adults who were mechanically ventilated. Both studies were single-centre RCTs. The interventions were similar, with Bucknall et al. (2008) indicating they modelled their intervention on that reported by Brook et al. (1999). In both studies, nurses used a structured approach for assessment to determine whether analgesics or sedatives (or both) were required by the patient, then
administered pre-specified medications according to their ongoing assessment. Differences in the medications used existed, with Brook et al. (1999) using diazepam, midazolam, fentanyl and morphine, while Bucknall et al. (2008) used midazolam, propofol and morphine.

The most important difference between the two studies was the usual method of providing sedation-related aspects of care to patients in each of the two study sites. In the US study, all aspects of sedation were ordered by the treating physicians and nurses could not make changes without a physician’s written or verbal order (Brook et al. 1999). In the Australian study, ICU medical staff prescribed the type of sedation medication and dose limits for infusion and boluses, with each patient’s ICU nurse free to assess, titrate and manage sedation, including the ceasing of sedation, in those limits (Bucknall et al. 2008).

Risk of bias in included studies

We analysed seven domains of potential risk of bias. We rated both studies the same for risk of bias for five of the seven domains (Table 2). Of note, usual care was not described well by Brook et al. (1999), except for the number of participants and duration of chemical paralysis. It was unclear if standard management practices (mode of mechanical ventilation, physiotherapy, suctioning, re-positioning, investigations outside ICU, need for physical restraints) or nurse:patient ratios were equally applied to both groups. While Bucknall et al. (2008) provided a description of usual care for general management and specific sedation management, some associated aspects of care, such as physiotherapy, suctioning, re-positioning, investigations outside ICU and need for physical restraints, were not provided. A potential for contamination between the two groups existed as participants in both studies were cared for in the same ICU at the same time and care of control group participants was directed by physicians in line with usual local practice and individual preferences (Brook et al. 1999, Bucknall et al. 2008). It is possible that the principles of protocol-directed care could have been partially applied to the control group.

Effects of interventions

Duration of mechanical ventilation

Both included studies reported duration of mechanical ventilation. When we pooled data to analyse the MD in duration of mechanical ventilation (MD = 5.74 hours, 95% CI 62.01 to 50.53) comparing management with protocol-directed sedation with usual care, the test of heterogeneity was substantial (τ² = 1416.10; χ² = 7.08, degrees of freedom (d.f.) = 1; P = 0.008; I² = 86%). Such high heterogeneity suggested that the two studies were dissimilar and
may reflect the differing nurse:patient ratios present in usual care in the study environments. Interpretation of these results related to duration of mechanical ventilation should proceed with caution given this high level of statistical heterogeneity.

**Intensive care unit and hospital mortality**

One study reported ICU mortality data (RR 1.04, 95% CI 0.67-1.61) (Bucknall et al. 2008) whereas both reported hospital mortality data. The combined hospital mortality outcome, with 633 patients, was not significantly different between the protocol-directed sedation and usual care groups (RR 0.96, 95% CI 0.71-1.31; heterogeneity Tau² = 0.02; \( \chi^2 = 1.50, \text{d.f.} = 1; P = 0.22; I^2 = 33\% \)) (Figure 2).

**Length of intensive care unit stay**

Both included studies reported length of ICU stay. The pooled data to analyse the MD in length of ICU stay (MD -0.62 days, 95% CI -2.97 to 1.73) comparing management with protocol-directed sedation with usual care, showed the test of heterogeneity was substantial (\( \chi^2 = 2.35; \chi^2 = 5.43, \text{d.f.} = 1; P = 0.02; I^2 = 82\% \)). Again, such high heterogeneity suggested that the two studies were dissimilar and interpretation of these results should proceed with caution.

**Hospital length of stay**

Both included studies reported hospital length of stay. The combined MD in hospital length of stay, with 633 patients, was not significantly different between the protocol-directed sedation and usual care groups (MD -3.78 days, 95% CI -8.54 to 0.97) (heterogeneity \( \chi^2 = 4.83; \chi^2 = 1.67, \text{d.f.} = 1; P = 0.20; I^2 = 40\% \)) (Figure 3).

**Adverse events**

The studies reported few adverse event data. One study reported re-intubation rates (RR 0.65, 95% CI 0.35-1.24) (Brook et al. 1999) while the other study reported self extubation data (RR 2.08, 95% CI 0.19-22.69) (Bucknall et al. 2008). In clinical practice, some patients who self extubate will not require re-intubation, therefore, self extubation rates would normally be higher than re-intubation rates. In these two studies, Bucknall et al. (2008) reported self extubation rates of only 1% in each group, while Brook et al. (1999) reported re-intubation rates of 6-13% in their two groups.

**Incidence of tracheostomy**

The incidence of tracheostomy was reported in both included studies. When we pooled data to analyse the frequency of tracheostomy (RR 0.77, 95% CI 0.31-1.89) comparing management with protocol-directed sedation with usual care, the test of heterogeneity was substantial (\( \chi^2 = 0.32; \chi^2 = 4.16, \text{d.f.} = 1; P = 0.04; I^2 = 76\% \)). Such high heterogeneity suggested that the two studies were dissimilar and interpretation of these results should proceed with caution. No studies were identified where the outcomes of total dose of sedation, incidence of delirium, memory function, psychological recovery, cognitive recovery or quality of life were addressed.

**Discussion**

**Summary of main results**

A systematic search of databases identified 2041 potential records, 21 potential studies and ultimately two eligible studies, with 633 participants, for review and analysis of the impact of protocol-directed sedation on duration of mechanical ventilation.

---

**Figure 2** Forest plot comparing protocol-directed sedation vs. non-protocol sedation to effect hospital mortality.
mechanical ventilation and mortality. Brook et al. (1999) reported a significant reduction in duration of mechanical ventilation and no difference in mortality with protocol-directed sedation in the US study, while Bucknall et al. (2008) reported no difference in either outcome in the Australian study. When we pooled data, hospital mortality did not differ between participants who received protocol-directed sedation and participants who received usual care. Significant heterogeneity suggested the cohorts were dissimilar for the outcome of duration of mechanical ventilation; therefore, interpretation of results should proceed with caution.

Secondary outcomes that were reported in both studies included ICU and hospital length of stay as well as incidence of tracheostomy. There was no difference in duration of hospital length of stay between participants who received protocol-directed sedation and participants who received usual care. Significant heterogeneity suggested the cohorts were very dissimilar for the outcomes of ICU length of stay and incidence of tracheostomy, therefore interpretation of results should proceed with caution.

Overall completeness and applicability of evidence

The two studies included in this systematic review both reported our primary outcomes; however, only a few of our secondary outcomes were reported. Neither study examined the relationship between protocol-directed sedation and post-ICU outcomes such as memory function, psychological and cognitive recovery and quality of life. Despite this, there is increasing recognition that sedation practices are likely to influence these long-term outcomes (Barr et al. 2013). Despite similar participant and intervention characteristics, substantial heterogeneity existed for most outcomes, limiting our ability to interpret the meta-analyses in a meaningful way. This heterogeneity may be the result of one study being conducted in the US in the 1990s (Brook et al. 1999), while the other study was conducted in Australia approximately 10 years later (Bucknall et al. 2008). These differences in geographical location and time may have resulted in substantial differences in important related areas of practice such as usual sedation practices and agents, patterns and modes of mechanical ventilation, mobilization practices and other aspects of intensive care that affect the identified outcomes. One aspect of critical care organization that differed between the two settings was the usual nurse:patient ratio, with each nurse caring for two or three patients in the US setting (confirmed with study investigators), while each nurse cared for one mechanically ventilated patient in the Australian setting; this has the potential to affect aspects of care such as how much patient agitation might be tolerated. Details regarding usual care are essential in the publication of studies that deal with a complex area of practice, as there are many variations that are essential to understand to determine transferability of evidence.

Quality of the evidence

The methodological quality of the studies was moderate, but the quality of the overall evidence was low. We only included two studies and they had conflicting results resulting in wide CIs for some outcomes. Furthermore, although we rated studies as having a low risk of detection and attrition bias and some aspects of selection bias, one or both studies had unclear or high risks of bias related to other

Figure 3 Forest plot comparing protocol-directed sedation vs. non-protocol sedation to effect hospital length of stay.
aspects of selection, reporting and performance. Due to the
nature of the intervention, it was not possible to blind par-
cipants or clinicians. Inclusion of alternative grades of evi-
dence, for example, non-randomized experimental studies
may help to provide a more complete picture of the evi-
dence, but is precluded under some Cochrane review group
guidelines. Furthermore, synthesis of qualitative studies
may be beneficial in identifying the characteristics of
patients and context, where nurse-directed sedation proto-
cols are beneficial and how benefit might be enhanced in
the future.

Potential biases in the review process
Clearly, described procedures were followed to minimize
potential bias in the review process. We conducted a sys-
tematic and rigorous literature search and used transparent
and reproducible methods. Where a review author was
involved in any included study, she was removed from the
process of analysing relevant information.

Agreements and disagreements with other studies or
reviews
The effect of the use of protocol-directed sedation on
patient outcomes has been of interest for several years
and, while it has not been the subject of any other
reviews, it has been the subject of additional, non-rando-
mized studies. Consistent with the findings of the two
studies included in this review (Brook et al. 1999, Bucknall
et al. 2008), findings from non-randomized studies have
generally been conflicting. One non-randomized study con-
ducted in Australia found no benefit and, in fact, an
increase in the duration of ICU length of stay with the
implementation of protocol-directed sedation (Elliott et al.
2006), while non-randomized studies conducted in Europe
identified mixed results. One Spanish study reported no
difference in duration of mechanical ventilation (Arias-Riv-
era et al. 2008), one Norwegian study reported a reduction
in duration of mechanical ventilation but no difference in
ICU length of stay (Brattebo et al. 2002) and two French
studies identified a reduction in duration of mechanical
ventilation (De Jonghe et al. 2005, Quenot et al. 2007).
These mixed results are likely to be influenced by multiple
behavioural factors in the study sites, particularly the role
of nurses in contributing to sedation management during
usual care.

One systematic review of observational and controlled
studies examined multiple aspects of sedation practice to
determine the impact of changes on economic and patient
safety outcomes (Jackson et al. 2010). When considering a
broad methodological range of studies, the overall conclu-
sion was that the introduction of guidelines and protocols
generally improved outcomes. Furthermore, in one related
systematic review of the effect of daily sedation interrup-
tion, there was no strong evidence of benefit from the inter-
vention although individual studies reported inconsistent
results (Burry et al. 2014). The reasons for these inconsis-
tencies are likely to be multidimensional; however, they
may include factors such as nurse:patient ratios, proportion
of speciality specific postgraduate educated nurses, sedative
agents used during usual care and other related aspects such
as ventilation and mobilization practices. It is also possible
that the sedation protocols resulted in different practices of
sedation administration that were not identified in the out-
comes assessed in this review. Both included studies mea-
sured doses of sedative agents but few differences were
noted and no total dose of sedation was available to enable
comparisons (Brook et al. 1999, Bucknall et al. 2008). It is
unlikely that any meaningful comparison of sedative agents
could be made given the effect of factors such as patient
weight and renal and liver function on drug metabolism.
Although inconsistencies in the effects of various interven-
tions have been identified, there is strong agreement that
the principle of reducing sedation, both in terms of depth
and duration, should be a goal of care given it is a link
with both short- and long-term outcome (Barr et al. 2013).
Achievement of this goal is likely to be optimized with con-
sistent use of validated assessment instruments, identifica-
tion of clear sedation targets and examination of various
interventions in local contexts.

Conclusion
Currently, limited evidence from RCTs is available to eval-
uate the effectiveness of protocol-directed sedation on
patient outcomes. The two included RCTs reported conflicting
results while heterogeneity limited the interpretation of
results for many of the outcomes. Notably, the clinical
context and practice roles of ICU clinicians should be con-
sidered prior to implementation of protocol-directed seda-
tion management. There was no evidence to draw
conclusions on the efficacy and safety of protocol-directed
sedation, although there was general agreement that vali-
dated sedation assessment instruments should be used in all
critical care settings and strategies to minimize sedation
should be implemented (Barr et al. 2013). The trend
towards sedation minimization has been ongoing since the
mid-2000s and is likely to continue, particularly in the
context of related strategies to optimize early mobilization.
and reduce complications of intensive care such as delirium and ongoing cognitive and psychological compromise (Needham et al. 2012).

Implications for research

Further research needs to be undertaken to ascertain the effect of protocol-directed sedation on patient outcomes. In particular, studies need to be conducted in a variety of clinical contexts to determine whether there are specific practice environments where benefit is more likely. The issue of whether a study randomized at the level of the individual can be conducted without contamination needs to be considered; it may be that a design such as cluster randomization is required. Given there are multiple different strategies that have been developed in recent years to reduce the detrimental impact of sedation, the interaction between protocol-directed sedation and other sedation minimization strategies should also be examined. It is vital that a detailed description of both the experimental care process and usual care is provided. Furthermore, a range of both process and outcome measures should be incorporated into the design, with outcome measures extending beyond confines of ICU or the acute care hospital and incorporating physical, cognitive and psychological health, and cost-effectiveness (Needham et al. 2012).

Acknowledgements

We thank Jane Cracknell (Managing Editor, Cochrane Anaesthesia Review Group) and Karen Hovhannisyan (Trials Search Co-ordinator, Cochrane Anaesthesia Review Group) for their assistance in the preparation of the protocol and review. We thank Harald Herkner (content editor), Nathan Pace (statistical editor) and John P Kress, Bronagh Blackwood and HS Jeffrey Mann (peer reviewers) and Janet Wale (CARG consumer editor) for their help and editorial advice during the preparation of this systematic review. We also thank Maria Isabel Castillo and Cesar Caramo for assistance with translating a Spanish language manuscript.

Funding

The National Health and Medical Research Council (NHMRC) has provided funding for this review from its Centre of Research Excellence scheme, which funds one or more of the authors. This research received no specific grant from any funding agency in the public, commercial or not-for-profit sectors.

Conflict of interest

No conflict of interest has been declared by the author(s).

Author contributions

All authors have agreed on the final version and meet at least one of the following criteria [recommended by the ICMJE (http://www.icmje.org/recommendations/)]:

- substantial contributions to conception and design, acquisition of data or analysis and interpretation of data;
- drafting the article or revising it critically for important intellectual content.

Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site.

References


The *Journal of Advanced Nursing (JAN)* is an international, peer-reviewed, scientific journal. *JAN* contributes to the advancement of evidence-based nursing, midwifery and health care by disseminating high quality research and scholarship of contemporary relevance and with potential to advance knowledge for practice, education, management or policy. *JAN* publishes research reviews, original research reports and methodological and theoretical papers.

For further information, please visit *JAN* on the Wiley Online Library website: www.wileyonlinelibrary.com/journal/jan

**Reasons to publish your work in JAN:**

- **High-impact forum:** the world’s most cited nursing journal, with an Impact Factor of 1.527 – ranked 14/101 in the 2012 ISI Journal Citation Reports © (Nursing (Social Science)).
- **Most read nursing journal in the world:** over 3 million articles downloaded online per year and accessible in over 10,000 libraries worldwide (including over 3,500 in developing countries with free or low cost access).
- **Fast and easy online submission:** online submission at http://mc.manuscriptcentral.com/jan.
- **Positive publishing experience:** rapid double-blind peer review with constructive feedback.
- **Rapid online publication in five weeks:** average time from final manuscript arriving in production to online publication.
- **Online Open:** the option to pay to make your article freely and openly accessible to non-subscribers upon publication on Wiley Online Library, as well as the option to deposit the article in your own or your funding agency’s preferred archive (e.g. PubMed).