Spontaneous Breathing Trials With T-Piece or Pressure Support Ventilation

José Augusto S Pellegrini MD, Rafael B Moraes PhD, Juçara G Maccari MD, Roselaine P de Oliveira PhD, Augusto Savi PhD, Rodrigo A Ribeiro PhD, Karen EA Burns PhD, and Cassiano Teixeira PhD

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Spontaneous breathing trials (SBTs) are among the most commonly employed techniques to facilitate weaning from mechanical ventilation. The preferred SBT technique, however, is still unclear. To clarify the preferable SBT (T-piece or pressure support ventilation [PSV]), we conducted this systematic review. We then searched the MEDLINE, EMBASE, SciELO, Google Scholar, CINAHL, ClinicalTrials.gov, and Cochrane CENTRAL databases through June 2015, without language restrictions. We included randomized controlled trials involving adult subjects being weaned from mechanical ventilation comparing T-piece with PSV and reporting (1) weaning failure, (2) re-intubation rate, (3) ICU mortality, or (4) weaning duration. Anticipating clinical heterogeneity among the included studies, we compared prespecified subgroups: (1) simple, difficult, or prolonged weaning and (2) subjects with COPD. We summarized the quality of evidence for intervention effects using the GRADE (Grading of Recommendations Assessment, Development, and Evaluation) methodology. We identified 3,674 potentially relevant studies and reviewed 23 papers in full. Twelve studies (2,161 subjects) met our inclusion criteria. Overall, the evidence was of very low to low quality. SBT technique did not influence weaning success (risk ratio 1.23 [0.94–1.61]), ICU mortality (risk ratio 1.11 [0.80–1.54]), or re-intubation rate (risk ratio 1.21 [0.90–1.63]). Prespecified subgroup analysis suggested that PSV might be superior to T-piece with regard to weaning success for simple-to-wean subjects (risk ratio 1.44 [1.11–1.86]). For the prolonged-weaning sub-
Introduction

Successful weaning of patients from mechanical ventilation constitutes one of the most challenging tasks for ICU practitioners. Timely identification of patients who are capable of spontaneous breathing can shorten the mechanical ventilation duration and potentially reduce mechanical ventilation-related complications.1-5

Once a patient is deemed ready to breathe spontaneously, a screening test, called a spontaneous breathing trial (SBT), is usually performed, although the literature remains conflicted on this subject.6-10 An SBT is typically performed by disconnecting the patient from the ventilator and attaching a T-piece to the endotracheal tube.11 Some clinicians, however, prefer to use low levels of pressure support ventilation (PSV), or automatic tube compensation.8

Switching from continuous mandatory ventilation to spontaneous breathing can decrease left-ventricular performance and unmask latent left-ventricular heart failure. Concerns exist regarding the potential for SBT failure rates to be higher with T-piece SBTs than with low levels of PSV, possibly because of the increased expenditure of respiratory muscle energy12 and cardiogenic pulmonary edema secondary to the Müller maneuver.13 Although PSV may be a less demanding SBT with regard to respiratory muscle effort and hydrostatic homeostasis, especially with the addition of PEEP to prevent the development of left-ventricular heart failure, it may also dull the clinical picture of intolerance compared with that of unassisted T-piece SBTs.13

Many trials have previously assessed this question, although heterogeneous methodological aspects and conflicting results limit adequate evidence appraisal. Previous meta-analyses have been conducted in this field but have not directly compared SBTs or used updated information.14

Objectives

Our objective was to clarify the preferred SBT technique (T-piece or low levels of PSV) for critically ill patients weaning from mechanical ventilation according to ICU mortality, reintubation rates, weaning failure, and weaning duration.

Methods

We conducted a systematic review based on standard methods and reported our findings in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement.15

Data Sources and Searches

We aimed to identify all randomized controlled trials (RCTs) assessing the efficacy and outcomes of T-piece compared with PSV trials in adult patients weaning from invasive mechanical ventilation.

We conducted electronic searches of the MEDLINE, EMBASE, Cochrane CENTRAL, CINAHL, SciELO, Google Scholar, and ClinicalTrials.gov databases for studies actively recruiting patients. All databases were searched from their inception to June 2015. Our MEDLINE search included the following terms and key words: (“weaning” OR “Ventilator Weaning”[MeSH]) AND (“Mechanical Ventilation” OR “Respiration, Artificial”[MeSH]) AND (“spontaneous breathing trial” OR “T-piece” OR “t tube”), using the Robinson and Dickersin RCT filter for PubMed.16 The electronic search strategy applied standard filters for the
identification of RCTs from each database. We screened the reference lists of retrieved publications for potentially eligible trials. We did not apply language restrictions.

Study Selection

We restricted our analysis to RCTs aiming to limit potential sources of bias. We excluded crossover trials and quasi-randomized trials. Regardless of specific weaning protocols, included trials had to compare between T-piece and PSV for conducting SBTs. We considered T-piece SBT to be the procedure of temporarily disconnecting a patient from the ventilator while maintaining an external oxygen supply, commonly by using a T-piece connected to the endotracheal tube. PSV was considered to be an SBT when employed in a systematic fashion, following a predefined protocol specifically designed to identify patients for extubation or, in the case of tracheostomized patients, for definitive removal from mechanical ventilation.

Outcome Measures

The outcomes assessed included (1) ICU mortality, (2) rate of re-intubation within 48 h following extubation, (3) weaning failure precluding extubation, and (4) weaning duration. We used authors’ definitions for the post-randomization weaning duration.

Data Extraction and Quality Assessment

Two independent reviewers (JP and RM) screened the titles and abstracts of retrieved citations and the full texts of potentially eligible studies to identify trials that met our inclusion criteria. Data from each potentially relevant trial were independently extracted by the reviewers using a predefined data extraction form.

According to the Cochrane risk of bias tool, we appraised the adequacy of random sequence generation, the reporting of allocation concealment, the blinding of participants and outcome assessments, and the descriptions of losses to follow-up and exclusions; we still assessed adherence to the intention-to-treat principle. We solved disagreements by consulting a third reviewer (CT) when needed.

Data Synthesis and Analysis

Qualitative Analysis. We used a narrative summary approach to qualitatively describe the study characteristics and variations in quality indicators and to consider how these factors affected our understanding of the outcomes of the included RCTs.

Quantitative Analysis. We used the Cochrane Collaboration guidelines to conduct our meta-analysis. All statistical analyses were performed with Review Manager version 5.3 (Nordic Cochrane Center, Copenhagen, Denmark), the Cochrane Collaboration’s software for preparing and updating Cochrane systematic reviews. We expressed the pooled effects estimates for binary and continuous variables using risk ratios and weighted mean differences with 95% CIs.

We tested for heterogeneity between studies using the Cochran Q and I² tests. We predefined statistical heterogeneity as being low, intermediate, or high, correlated to I² statistics <25%, from 25 to 50%, or >50%, respectively. Meta-analyses with random-effects models were employed for all outcomes, due to anticipated clinical heterogeneity in terms of patient populations. We attempted to identify clinical factors as potential sources of heterogeneity, assessing for prespecified subgroups, including (1) weaning difficulty (simple, difficult, or prolonged weaning) and (2) COPD (vs subjects without COPD).

To assess the potential publication bias from small study effects, we constructed funnel plots displaying the log risk ratio on the horizontal axis and the SE of the log risk ratio on the vertical axis. We employed Egger’s test to evaluate the risk for publication bias. We summarized the quality of evidence according to the Grading of Recommendations Assessment, Development, and Evaluation (GRADE) guidelines and reported them using the GRADE pro web-based tool.

Results

Study Selection

Our initial electronic search identified 3,674 abstracts. Of these, we excluded 3,651 because they did not describe RCTs, did not evaluate weaning techniques, were duplicate references, or were not relevant. We retrieved 23 studies for a more detailed, full-text analysis, and we excluded 11 of these studies (Fig. 1). We then identified 12 suitable studies comprising 2,161 subjects. Both reviewers completely agreed on the final selection of included studies. We also identified 2 ongoing RCTs, including one from our group, from the ClinicalTrials.gov database (Pellegrini, http://clinicaltrials.gov/show/NCT01464567; Agarwal, http://clinicaltrials.gov/show/NCT00911378 Accessed August 15, 2016).

Study Description

The included studies were published between 1994 and 2015 and were from 10 countries. Eight of the 12 included studies were single-center studies. One study was pub-
lished only in abstract form,37 and full details of the study were not available to the authors of this review.

Table 1 summarizes the components of the risk of bias assessment. Due to the nature of the intervention being studied, all studies were unblinded with regard to the patients enrolled and the outcomes assessed. Only 3 studies specified adequate random sequence generation, and 7 studies did not report adherence to the intention-to-treat principle. Our qualitative analysis of key study characteristics is summarized in Table 2.

Outcomes Assessed

All but 2 trials31 included in the study reported weaning failure outcome, which was defined as failure to extubate the subject immediately following the SBT. All but one36 study, apart from those assessing only tracheostomized subjects, reported 48-h re-intubation rates. Eight studies reported ICU mortality.32-35,37-39,41

Study Protocols

Brochard et al38 compared 3 different strategies for gradual weaning from mechanical ventilation. Beyond T-piece and PSV, the authors randomly assigned subjects to a third group using gradual titration of synchronized intermittent mandatory ventilation. T-piece SBTs were performed up to 8 times/d, progressing from 5 to 120 min in progressive steps. In subjects assigned to the PSV group, the PSV level was systematically adjusted 2 times/d to maintain a breathing frequency between 20 and 30 breaths/min in decremental steps of 2–4 cm H2O. Esteban et al40 compared once-a-day T-piece SBTs with 3 other methods, including intermittent mandatory ventilation, intermittent trials of spontaneous breathing (conducted two of more times a day if possible), in addition to PSV.

The study by Vitacca et al41 differed from the other included studies by enrolling only difficult-to-wean tracheostomized subjects with COPD who required mechanical ventilation for at least 15 d. This trial was conducted in 3 long-term weaning units, with subjects transferred from 24 ICUs after a range of 15–39 d on mechanical ventilation. The authors also compared their results with historical controls or an uncontrolled clinical practice. Jubran et al34 assessed a similar population. Subjects who required mechanical ventilation for >21 d in a long-term weaning unit were randomly assigned to unassisted breathing through a tracheostomy collar or to progressive reductions in PSV based on their breathing frequencies.

Six studies included simple-to-wean subjects, according to previously published definitions.2,30-32,35,36,39 Three studies assessed difficult-to-wean subjects,33,38,40 and 2 studies included prolonged-weaning and tracheostomized subjects.34,41 Two additional studies evaluated postoperative subjects.35,37

In the studies that predefined the SBT duration, a 120-min trial was most commonly employed (7 studies). Three authors reported progressively increasing duration of SBTs based on subjects’ tolerance.34,38,41

Although included in various trials, subjects with COPD represented only a small fraction of the study population. Only 2 of the included studies specifically assessed subjects with COPD33,41; one of these also specifically enrolled prolonged-weaning subjects.41

Evidence Synthesis

T-piece SBTs were associated with a risk ratio (95% CI) of 1.11 (0.80–1.54) for ICU mortality and 1.21 (0.90–1.63) for the 48-h re-intubation rate (Figs. 2 and 3). The evidence from trials addressing these outcomes was considered very low to low quality based on the GRADE approach (Table 3). Study limitations, inconsistency, and imprecision contributed to downgrading the overall quality of evidence in the pooled RCTs.

For weaning failure, we found a risk ratio of 1.23 (0.94–1.61) (Fig. 4) with moderate to high heterogeneity (I2 = 48%). When evaluating potential sources of clinical heterogeneity, we excluded prolonged-weaning studies from our analysis (8 studies remaining; 1,237 subjects) and noted a risk ratio of 1.47 (1.17–1.84) favoring PSV with regard to weaning success. The I2 statistic for this analysis was 0%, suggesting that prolonged-weaning studies represent an important source of clinical heterogeneity.
**SBTs With T-Piece or PSV**

We performed funnel-plot analysis for each outcome and did not identify publication bias. The funnel plot for weaning failure is shown in Figure 6. Egger’s test did not suggest publication bias ($P = .37$).

### Discussion

**Summary of Evidence**

T-piece and PSV are 2 of the most commonly used techniques when conducting SBTs in clinical practice today. Nevertheless, existing evidence directly comparing these 2 approaches is sparse, heterogeneous, and of poor overall quality. Small study populations with low event rates, variability among the applied SBT techniques, and remarkably different populations limit the pooling and adequate interpretation of evidence.

T-piece and PSV techniques have theoretical singularities that may influence bedside judgment when choosing one SBT technique over another. When using T-piece SBTs, one might be looking for specificity and thus might proceed with extubation only for those patients able to tolerate the hemodynamic perturbations of this disturbing test.\(^{13,42}\) Also, previous studies have shown that the post-extubation work of breathing could be more closely paralleled by unassisted breathing (as in a T-piece trial) than by a low-pressure support trial.\(^{43}\)

By contrast, Ezingeard et al\(^ {44}\) demonstrated that more subjects could be successfully extubated after a PSV SBT, including some subjects who previously failed a T-piece SBT. These findings are supported by moderate-quality evidence with regard to simple-to-wean subjects, for whom PSV might be associated with reduced weaning failure rates, not adversely influencing re-intubation rates.

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### Table 1. Risk of Bias Assessment

<table>
<thead>
<tr>
<th>Study</th>
<th>Adequate Sequence Generation</th>
<th>Allocation Concealment</th>
<th>Blinding of Participants and Investigators</th>
<th>Blinding of Outcome Assessment</th>
<th>Description of Losses and Exclusions</th>
<th>Intention to Treat Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brochard et al(^ {38})</td>
<td>Not clear</td>
<td>SNOSE</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Esteban et al(^ {39})</td>
<td>Yes</td>
<td>SNOSE</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Not clear</td>
</tr>
<tr>
<td>Esteban et al(^ {40})</td>
<td>Yes</td>
<td>SNOSE</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Not clear</td>
</tr>
<tr>
<td>Vitacca et al(^ {41})</td>
<td>Not clear</td>
<td>Not clear</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Not clear</td>
</tr>
<tr>
<td>Haberthur et al(^ {42})</td>
<td>Not clear</td>
<td>SNOSE</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Koksal et al(^ {43})</td>
<td>Not clear</td>
<td>Not clear</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Matič et al(^ {44})</td>
<td>Not clear</td>
<td>SNOSE</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Not clear</td>
</tr>
<tr>
<td>Matič et al(^ {45})</td>
<td>Not clear</td>
<td>SNOSE</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Not clear</td>
</tr>
<tr>
<td>Zubran et al(^ {46})</td>
<td>Not clear</td>
<td>SNOSE</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Lourenço et al(^ {47})</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Not clear</td>
</tr>
<tr>
<td>Zhang et al(^ {48})</td>
<td>Not clear</td>
<td>Not clear</td>
<td>No</td>
<td>No</td>
<td>Not clear</td>
<td>Not clear</td>
</tr>
<tr>
<td>Chittawatanarat et al(^ {49})</td>
<td>Not clear</td>
<td>Not clear</td>
<td>No</td>
<td>No</td>
<td>Not clear</td>
<td>Not clear</td>
</tr>
</tbody>
</table>

SNOSE = sequentially numbered, opaque, sealed envelopes

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### Subgroup Analyses

**Weaning Difficulty.** Seven studies (1,600 subjects) focused on simple weaning, which was defined as success on the first SBT in the absence of previous weaning failure. PSV in this population was associated with better outcomes related to weaning success (risk ratio = 1.44, 1.11–1.86; I\(^2\) = 0%) but not with lower re-intubation or ICU mortality rates. Three studies (197 subjects) specifically assessed difficult-to-wean subjects. In this subgroup, significant differences in clinical outcomes were not found between the alternative SBT techniques.

Only 2 studies, comprising 364 subjects, focused on prolonged-weaning subjects. Whereas the SBT technique had no effect on weaning failure rate or ICU mortality, T-piece was associated with a shorter weaning duration (weighted mean difference = −3.08 (−5.24 to −0.92) d of weaning) than that associated with PSV (Fig. 5). These aforementioned outcomes had an I\(^2\) = 0% in this subgroup.

**COPD Subjects.** In the 12 included studies, we identified 338 subjects with COPD, although 3 studies did not specifically report this condition.

Between the 2 studies enrolling exclusively subjects with COPD, we identified remarkable clinical heterogeneity preventing pooling. Whereas the study of Matič et al\(^ {43}\) included difficult-to-wean subjects with COPD (defined as one failed weaning attempt), the Vitacca study\(^ {41}\) included tracheostomized subjects ventilated for $\geq$15 d. The risk ratios in the Matič and Vitacca studies for weaning failure with T-piece SBTs were 1.61 (0.82–3.16) and 0.86 (0.33–2.21), respectively, and risk ratios for ICU mortality were 2.06 (0.41–10.47) and 0.67 (0.12–3.67), respectively.
Table 2. Qualitative Analysis of Key Study Characteristics

<table>
<thead>
<tr>
<th>Study</th>
<th>COPD</th>
<th>SBT Duration (min)</th>
<th>Mode of Weaning</th>
<th>Pre-SBT (Mean ± SD)</th>
<th>Days on Mechanical Ventilation</th>
<th>Re-intubation Within 48 h</th>
<th>Weaning Failing</th>
<th>Mortality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brochard et al.44</td>
<td>Yes</td>
<td>Simple</td>
<td>120</td>
<td>Yes</td>
<td>4.6 ± 3.4</td>
<td>120</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Esteban et al.45</td>
<td>Yes</td>
<td>Difficult</td>
<td>24</td>
<td>Yes</td>
<td>4.7 ± 3.5</td>
<td>47</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Vitacca et al.46</td>
<td>Yes</td>
<td>Simple</td>
<td>120</td>
<td>Yes</td>
<td>4.6 ± 3.5</td>
<td>47</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Haberthür et al.30</td>
<td>No</td>
<td>Simple</td>
<td>120</td>
<td>Yes</td>
<td>4.6 ± 3.5</td>
<td>100</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Maticetti et al.33</td>
<td>No</td>
<td>Simple, post-surgical</td>
<td>120</td>
<td>Yes</td>
<td>4.6 ± 3.5</td>
<td>NR</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Zhang et al.36</td>
<td>No</td>
<td>Difficult</td>
<td>120</td>
<td>Yes</td>
<td>4.6 ± 3.5</td>
<td>100</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Chittawatanarat37</td>
<td>No</td>
<td>Simple, post-surgical</td>
<td>NR</td>
<td>No</td>
<td>4.6 ± 3.5</td>
<td>NR</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

In contrast, low-quality evidence suggests that prolonged-weaning subjects\(^31,\)\(^57\) appear to benefit from T-piece SBTs in terms of weaning duration. In these subjects, progressive steps toward predetermined reductions in PSV according to the subject’s tolerance may prolong the duration of mechanical ventilation, potentially increasing the risk for mechanical ventilation-related complications. However, we observed that the few studies including subjects who experienced such events found that the SBT technique has no influence on mortality in this subgroup.

Patients with COPD represent a growing population worldwide, remaining as one of the most prevalent conditions leading to respiratory failure and consequent mechanical ventilation.\(^45-47\) These individuals represent some of the most challenging groups to wean from mechanical ventilation\(^48;\) paradoxically, this population is underrepresented in RCTs. Two groups of authors evaluated subjects with COPD exclusively, but their studies included markedly different profiles. One group of authors\(^33\) enrolled difficult-to-wean subjects with COPD, and the other\(^41\) focused on tracheostomized, prolonged-weaning subjects in long-term weaning units. Recognizing that clinical heterogeneity would hinder the interpretation of findings, we decided not to pool these results.

A relevant aspect that should be kept in mind concerns the consideration of SBTs as an intervention rather than a diagnostic test trying to identify patients who are potential candidates for extubation, predicting tolerance of unassisted breathing. In the latter approach, questions focus on diagnostic accuracy, and then weaning failure or re-intubation rates are important end points for describing weaning trial sensitivity or specificity.

Nevertheless, the assessment of diagnostic properties of SBTs for predicting successful extubation is not straightforward. Extubation failure rates are widely reported as being around 15–20%, which makes specificity of the trial for predicting successful extubation only 80–85%. On the opposite side, test sensitivity (the proportion of patients able to tolerate extubation despite failing the weaning test) is difficult to evaluate because patients who fail a weaning test are usually not extubated. Furthermore, criteria for termination of a weaning trial and even definitions of test failure are essentially subjective and clinician-dependent, potentially biasing outcomes beyond the test itself. Accordingly, we decided to stay in line with previous studies and assess the clinical impact of SBTs as an intervention in important outcomes, beyond its diagnostic role in predicting patient tolerance to mechanical ventilation discontinuation.

Overall, our results are consistent with those of a Cochrane review\(^14\) and are applicable to the general weaning population encountered by clinicians in clinical practice. Our review, however, adds important additional information from 4 published RCTs, increasing the size of the included population (2,161 subjects here vs 1,208 subjects...
previously analyzed). In addition, we defined an a priori subgroup analysis aimed at identifying different effects of the alternative SBT techniques based on weaning difficulty and the presence of COPD. Very low to low overall quality of evidence strongly limits definitive findings in this field.

**Strengths and Limitations**

We conducted a systematic search of several databases without language restrictions to identify all RCTs comparing T-piece and PSV SBT techniques in weaning subjects. We employed standardized techniques to assess risk of bias and overall quality of evidence.

Our review has several limitations that reduce the strength of inferences that can be made. First, quality assessment permits classifying the evidence as very low to low quality. Very low to low quality of evidence strongly limits definitive findings in this field.

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**Fig. 2.** ICU mortality. M-H = Maentel-Haentzel; PSV = pressure support ventilation.

**Fig. 3.** 48-h reintubation rate. M-H = Maentel-Haentzel; PSV = pressure support ventilation.
Table 3. Grade Evidence Profile

<table>
<thead>
<tr>
<th>Study Event Rates (%)</th>
<th>Relative effect (95% CI)†</th>
<th>Anticipated Absolute Effects on Study Population</th>
</tr>
</thead>
<tbody>
<tr>
<td>With PSV</td>
<td>With T-Piece</td>
<td>Risk With PSV</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Risk Difference With T-Piece</td>
</tr>
<tr>
<td>ICU mortality: simple weaning</td>
<td>62/903 (6.9) RR 1.11 (0.80 to 1.54)</td>
<td>691,000 8 more/1,000 (14 fewer to 37 more)</td>
</tr>
<tr>
<td>ICU mortality: difficult weaning</td>
<td>31/662 (4.7) RR 1.32 (0.84 to 2.07)</td>
<td>471,000 15 more/1,000 (7 fewer to 50 more)</td>
</tr>
<tr>
<td>ICU mortality: prolonged weaning</td>
<td>6/63 (9.5) RR 1.86 (0.75 to 4.62)</td>
<td>951,000 82 more/1,000 (24 fewer to 345 more)</td>
</tr>
<tr>
<td>ICU mortality: COPD exclusive</td>
<td>140/1,000 RR 0.96 (0.61 to 1.51)</td>
<td>140,000 15 more/1,000 (24 fewer to 345 more)</td>
</tr>
<tr>
<td>Re-intubation rate: simple weaning</td>
<td>75/712 (10.5) RR 1.24 (0.79 to 2.01)</td>
<td>1051,000 31 per 1,000 (12 fewer to 117 more)</td>
</tr>
<tr>
<td>Re-intubation rate: difficult weaning</td>
<td>16/100 (16.0) RR 1.21 (0.66 to 2.22)</td>
<td>161,000 34 more/1,000 (54 fewer to 195 more)</td>
</tr>
<tr>
<td>Re-intubation rate: prolonged weaning</td>
<td>194/903 (24.2) RR 1.23 (0.94 to 1.61)</td>
<td>2421,000 56 more/1,000 (14 fewer to 147 more)</td>
</tr>
<tr>
<td>Duration of weaning</td>
<td>83/525 (15.8) RR 1.44 (1.11 to 1.86)</td>
<td>158,000 70 more/1,000 (17 more to 136 more)</td>
</tr>
<tr>
<td>ICU mortality: COPD exclusive</td>
<td>91/187 (51.1%) RR 0.85 (0.69 to 1.05)</td>
<td>511,000 77 fewer/1,000 (158 fewer to 26 more)</td>
</tr>
<tr>
<td>ICU mortality: COPD exclusive</td>
<td>16/58 (27.6%) RR 1.28 (0.71 to 2.32)</td>
<td>276,000 77 more/1,000 (80 fewer to 364 more)</td>
</tr>
</tbody>
</table>

* Unclear sequence generation in 9 studies as well as allocation concealment in 5 studies; patients or outcome assessors were not blinded in any study; intention to treat analysis in only 5 studies.
† Relative effects (95% CI) are based on random-effect models.
RCT = randomized controlled trial
RR = relative risk
MD = mean difference
methodological issues that may impact study findings. Im-
precision of available data was an important source of
downgrading of evidence for many outcomes. Second, we
identified important clinical heterogeneity among studies
that hindered the pooling of estimates and limited the gen-
eralizability of our findings. Some aspects related to me-
chanical ventilation settings (different pressure support lev-
els and protocols, adjunctive use of PEEP) certainly
contribute to conflicting results. This could be considered one
of the most important issues in this review. Third, subgroup
analysis should be interpreted with caution, accordingly to
study populations and outcomes reported. Finally, the diffi-
ticult-to-wean subject and COPD subject subgroups remain
scarcely studied, limiting conclusions in these areas.

Conclusions

The quality of available evidence precludes definitive
conclusions about assessed outcomes. Low-quality evi-
dence suggests that PSV SBTs may result in lower wean-
ing failure rates in simple-to-wean subjects but do not
affect re-intubation rates or other important outcomes. Con-
versely, in prolonged-weaning subjects, a T-piece may re-
duce the weaning duration compared with PSV SBTs.

Fig. 4. Weaning failure. M-H = Maentel-Haentzel; PSV = pressure support ventilation.

Fig. 5. Forest plot comparing T-piece and pressure support ventilation (PSV) for weaning duration. M-H = Maentel-Haentzel.

Fig. 6. Funnel plot for weaning failure outcome.
SBTs With T-Piece or PSV

Future trials should compare SBT techniques in difficult-to-wean and COPD subjects.

REFERENCES


