Effectiveness of bronchoscopic lung volume reduction using unilateral endobronchial valve: a systematic review and meta-analysis

Background: Bronchoscopic lung volume reduction (BLVR) can be suggested as an alternative for surgical lung volume reduction surgery for severe emphysema patients. This article intends to evaluate by systematic review the safety and effectiveness of BLVR using a one-way endobronchial valve.

Methods: A systematic search of electronic databases, including MEDLINE, EMBASE, and the Cochrane Library, as well as eight domestic databases up to December 2013, was performed. Two reviewers independently screened all references according to selection criteria. The Scottish Intercollegiate Guidelines Network criterion was used to assess quality of literature. Data from randomized controlled trials were combined and meta-analysis was performed.

Results: This review included 15 studies. Forced expiratory volume in 1 second (FEV1) improved in the intervention group compared with the control group (mean difference [MD]=6.71, 95% confidence interval [CI]: 3.31–10.11). Six-minute walking distance (MD=15.66, 95% CI: 1.69–29.64) and cycle workload (MD=4.43, 95% CI: 1.80–7.07) also improved. In addition, St George’s Respiratory Questionnaire score decreased (MD=4.29, 95% CI: –6.87 to –1.71) in the intervention group. In a subgroup analysis of patients with complete fissure, the FEV1 change from baseline was higher in the BLVR group than in the control group for both 6 months (MD=15.28, P<0.001) and 12 months (MD=17.65, P<0.001), whereas for patients with incomplete fissure, FEV1 and 6-minute walking distance showed no change. One-year follow-up randomized controlled trials reported deaths, although the cause of death was not related to BLVR. Respiratory failure and pneumothorax incidence rates were relatively higher in the BLVR group, but the difference was not significant.

Conclusion: BLVR may be an effective and safe procedure for the treatment of severe COPD patients with emphysema, based on existing studies.

Keywords: endobronchial valve, bronchoscopic valve lung volume reduction, systematic review, meta-analysis

Introduction

Chronic obstructive pulmonary disease (COPD) is a respiratory disease characterized by irreversible abnormal lung inflammation and shows progressive airflow limitation.1 It is the fifth leading cause of death in high-income countries2 and is one of the major public health problems across the world. According to the Global initiative for chronic Obstructive Lung Disease (GOLD), it is predicted that COPD will be the third cause of death in 2020.3 To reduce hyperinflation, lung volume reduction surgery for severe emphysema has been performed for over 30 years. Surgery-related complications and mortality are the main concerns with this form of treatment.4
Bronchoscopic lung volume reduction (BLVR) was introduced to maintain the effect of lung volume reduction and also reduce surgery-related complications and mortality. Many studies of BLVR have been published, with the majority of study designs being case series. Therefore, randomized controlled trials (RCTs) have been recommended for evidence enhancement. Recently, the US and Europe have reported results of multicenter RCTs using unilateral bronchial valves for BLVR, but meta-analysis is not yet published. This study is a systematic review to evaluate the effectiveness and safety of BLVR using unilateral bronchial valves in COPD with emphysema.

Materials and methods

Search strategy

A literature search was performed using eight domestic research databases, including KoreaMed, and core databases such as MEDLINE, EMBASE, and the Cochrane Library up to December 2013. Key question and patient intervention comparators and outcomes were defined under the advice of expert groups. The presearch included 148 article abstracts. Extensive searches of databases using the terms “endoscopic lung volume reduction”, “bronchoscopic lung volume reduction”, “endobronchial valve”, and “BLVR” were performed. Terms were related to the intervention and modified according to each database’s index term, such as Medical Subject Heading (MeSH) and EMTREE.

Inclusion/exclusion criteria

Studies that met the following criteria were included: 1) study population is people with COPD with severe emphysema; 2) intervention is BLVR using a unilateral bronchial valve; 3) study design is an RCT, a cohort study, or a case series; 4) reported one of the predetermined outcomes; and 5) published in Korean or English.

Studies on animal trial or preclinical studies and nonoriginal articles such as reviews, editorial letters, and comments were excluded. Articles not published in either Korean or English and studies with duplicate subjects (study using the same outcome indicators published in duplicate) were also excluded.

Types of outcome measures

Primary outcome was lung function at forced expiratory volume in 1 second (FEV1). Secondary outcomes were 6-minute walk distance (6MWD), cycle workload (Watt), quality of life: St George’s Respiratory Questionnaire (SGRQ) score, dyspnea scale: modified Medical Research Council score, and safety issues: major complications (death, pneumothorax, massive hemoptysis) and minor complications (pneumonia, minor hemoptysis, COPD exacerbation).

Data collection and analysis

Selection of studies and quality assessment

Each stage, from the literature search to the application of selection criteria and data extraction, was independently undertaken by two researchers. Study selection was carried out by predefined inclusion/exclusion criteria. An expert group consisting of respiratory medicine and evidence-based medicine specialists and thoracic surgery specialists in COPD guided each stage. Quality assessment of selected studies was carried out using tools of the Scottish Intercollegiate Guidelines Network.

Data extraction and management

According to a preagreed data extraction format, two investigators independently extracted data for review. Continuous variables such as mean change from baseline, median range, and standard deviation were converted according to the formula in the Cochrane Handbook for Systematic Reviews of Interventions. The extracted data syntheses were performed both quantitatively and qualitatively.

Measurement and treatment effect

The mean difference (MD) and 95% confidence interval (CI) from each individual RCT were calculated, and meta-analysis using a fixed effect model was performed. Two RCTs were included in the meta-analysis. One set of data from an RCT was not included, because the comparator and follow-up period were different. Statistical analysis was performed using Cochrane RevMan version 5.2 and Comprehensive Meta Analysis software.

Results

Study characteristics

A total of 1,016 studies were retrieved from the database. After exclusion of duplicates, 678 studies remained. Finally, 15 studies were selected according to the selection and exclusion criteria (Figure 1): three RCTs, one cohort study, and eleven case studies. All studies used a duckbill-type endobronchial valve. Six studies used a Zephyr valve and eight studies used an Emphasis® valve (Table 1).
Identified studies through electronic searches of databases (total n=1,016)
- Ovid-MEDLINE (n=329), Ovid-EMBASE (n=633)
- Cochrane library (n=46)
- Domestic databases and hand-search (n=8)

Remaining studies after excluding duplicates (n=678)
Full-text articles assessed for eligibility (n=232)
Studies included in synthesis (n=15)

Excluded by selection criteria (n=446)
1. Animals or pre-clinical studies (n=22)
2. Not original articles (n=230)
3. Gray literatures (n=95)
4. Case reports (n=53)
5. Did not published in English or Korean (n=46)

Excluded studies (n=217)
1. Patients did not have COPD with emphysema (n=68)
2. Not performed BLVR intervention (n=123)
3. Did not report outcomes of our interest (n=20)
4. Patient duplicates as same outcomes (n=6)

Clinical effectiveness
Two RCTs included in the meta-analysis compared the intervention group with the control group that performed only medical treatment. Each outcome measure, such as FEV₁, 6MWD, cycle workload, and SGRQ, was reported (Figure 2). Six months after, the mean change from baseline and standard error were calculated for each group. The MDs with 95% CI were compared between the intervention group and the control group. The FEV₁ improved in the intervention group compared with the control group (MD=6.71, P<0.0001). 6MWD and cycle workload in the intervention group also increased (MD=15.66, MD=4.43). The SGRQ score in the intervention group decreased compared with the control group (MD=4.29).

Although not included in the meta-analysis, the other RCT reported that for BLVR compared with the sham procedure (bronchoscopy) over 3 months, computed tomography (CT) lung volume and number of SGRQ responders were increased in the intervention group by 24.2% (8/33) compared with the control group (0/33). However, FEV₁ and 6MWD were no different between the two groups.

One retrospective cohort study performed quantitative CT image analysis before and after the procedure. After 6 months of observation of lung volume, target lobe volume in the intervention group decreased from baseline (−0.451 cc, P<0.0001), but not in the control group (0.0051 cc, P=0.70).

Clinical effectiveness according to fissure completeness
The subgroup was evaluated for the presence of complete fissure, and several outcomes were compared. The results are shown in Table 2. For patients with complete fissure, the FEV₁ change from baseline in the BLVR group was higher than in the control group for both 6 months (MD=15.29, P<0.001) and 12 months (MD=17.65, P<0.001). However, for the patients with incomplete fissure, mean change of FEV₁ and 6MWD was not different.
In addition, one case series study reported that 5-year survival rate of the fissure group was 83.3%, whereas the nonfissure group was 24.0% by log-rank test ($P=0.0036$).

### Safety-related outcomes

Three RCTs reported safety,\textsuperscript{7,8,12} two of them\textsuperscript{7,8} compared BLVR with medical treatment, and the other RCT\textsuperscript{12} compared BLVR with sham bronchoscopy. Firstly, one of the RCTs\textsuperscript{7} compared BLVR with medical treatment and found that mortality over 12 months occurred in 5.4% of the BLVR group of 111 participants and in 6.7% of the control group of 60 participants, although it was not statistically different. An independent committee of the RCT judged that the death and the BLVR procedure were not related. Respiratory failure occurred only in the intervention group (9.0%). However, after 12 months, the occurrence of respiratory failure was not different between the two groups. The incidence of pneumothorax after the procedure occurred only in the intervention group (8.2%), yet the incidence of pneumothorax was not statistically different. Massive hemoptysis occurred only in the intervention group (0.9%).

In the other RCT study,\textsuperscript{8} death occurred only in the intervention group (2.7%) of 220 participants, although there was no significant difference between the two groups ($P=0.19$). Causes of death were three cases of respiratory failure not associated with BLVR, one ischemic colitis, one cancer, and one massive hemoptysis. Respiratory failure occurred only in the intervention group (1.4%). Pneumothorax occurred only in the BLVR group (4.1%). One massive hemoptysis occurred in the intervention group. There was no statistically significant difference between the BLVR group and the control group.

One RCT\textsuperscript{12} compared BLVR with bronchoscopy as a sham procedure. During the 3-month follow-up period, adverse events were classified according to severity grade: serious, severe, moderate, or mild. Serious adverse events were seen in the intervention group in 18.9% of 37 participants and in 11.1% of the control group of 36 participants. The incidence rate was not different between the two groups ($P=0.214$).

Because the follow-up period of RCT studies was less than 12 months, ten case series studies were selected for safety review. The follow-up observation period ranged from 1 month up to 5 years. Three of the studies\textsuperscript{19,21,23} reported death, but it was not related to the procedure. Of the major complications, seven pneumothorax cases were reported, and the incidence rate ranged from 2.5% (2/40) to 23% (3/13). Pneumonia distal to valve was 5.2% (1/19) in one study.\textsuperscript{18}


Discussion

BLVR using a one-way valve is a procedure for COPD patients with severe emphysema. Theoretically, a one-way valve inserted into the bronchus helps air outflow from the target lobe during expiration, and blocks air inflow into the lobe during inspiration. This mechanism improves the overall lung function and relieves the symptom of dyspnea by reducing the lung volume of the target lobe. However, even after the procedure, there are often collapse failures by collateral ventilation, in which case the effect of the procedure is reduced. The known appropriate population for the BLVR procedure is heterogeneous emphysema patients, and currently two types of bronchial valve are used. One is the duckbill-type valve; the other is the umbrella valve. Differences in the structure of these valves may impact on efficacy and safety, so only the duckbill-type valve was considered in this study.

The UK and European health technology assessment reports, based on a synthesis of the literature, did not include RCT studies and focused on case studies. For evaluation of safety and efficacy related to BLVR, these assessments recommended more RCTs. In a large RCT study, the Endobronchial Valve for Emphysema Palliation Trial (VENT) was included in the published results. VENT’s selection criteria were similar to those for surgical lung volume reduction. Only 6 months of data were
Table 2 Clinical outcomes in change from baseline according to fissure

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Author (year)</th>
<th>Quality</th>
<th>N</th>
<th>Percent change difference between BLVR vs control</th>
<th>6 months</th>
<th>12 months</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>BLVR</td>
<td>Control</td>
<td>MD (95% CI)</td>
</tr>
<tr>
<td>Complete fissure</td>
<td>Herth et al (2012)</td>
<td>I+</td>
<td>44</td>
<td>14 (5.16 to 22.84)</td>
<td>4.51</td>
<td>15.29 (P&lt;0.001)</td>
</tr>
<tr>
<td></td>
<td>Sciruba et al (2010)</td>
<td>I+</td>
<td>68</td>
<td>16.2 (8.80 to 23.50)</td>
<td>3.75</td>
<td>17.9 (P=0.025)</td>
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<td></td>
<td></td>
<td></td>
<td>33</td>
<td>8 (3.43 to 18.28)</td>
<td>13.41</td>
<td>3 (P=0.192)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>32</td>
<td>7.7 (-1.80 to 17.20)</td>
<td>4.85</td>
<td>3.9 (-4 to 11.8)</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>19</td>
<td>7.0 (1.80 to 12.20)</td>
<td>2.65</td>
<td>6 (0.21 to 11.79)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>19</td>
<td>-9 (-17.07 to -0.93)</td>
<td>4.12</td>
<td>-4 (-10.64 to -2.65)</td>
</tr>
<tr>
<td>Incomplete fissure</td>
<td>Herth et al (2012)</td>
<td>I+</td>
<td>67</td>
<td>2 (-5.81 to 9.81)</td>
<td>3.98</td>
<td>2 (-6.06 to 10.06)</td>
</tr>
<tr>
<td></td>
<td>Sciruba et al (2010)</td>
<td>I+</td>
<td>29</td>
<td>2 (-3.9 to 7.9)</td>
<td>3.01</td>
<td>2.8 (-3.8 to 9.4)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>56</td>
<td>5 (-5.44 to 15.44)</td>
<td>5.33</td>
<td>5 (-2.73 to 12.73)</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>56</td>
<td>3 (-1.5 to 12.2)</td>
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<td>4.5 (-2.7 to 11.8)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>40</td>
<td>3 (-1.7 to 7.78)</td>
<td>2.44</td>
<td>4 (-1.09 to 9.09)</td>
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<td></td>
<td></td>
<td></td>
<td>40</td>
<td>-3 (-8.48 to 2.80)</td>
<td>2.80</td>
<td>0 (-5.48 to 5.48)</td>
</tr>
</tbody>
</table>

Abbreviations: BLVR, bronchoscopic lung volume reduction; MD, mean difference; CI, confidence interval; SE, standard error; ΔFEV1, change in forced expiratory volume in 1 second; ΔMWD, change in 6-minute walking distance; ΔCycle workload, change in cycle workload; ΔSGRQ, change in St George's Respiratory Questionnaire.
of complications was not different between the intervention group and the control group. Most patients involved in these studies had severe emphysema and were considered a high-risk population for receiving surgery. Since BLVR is a less invasive procedure than surgery, safety could be determined as acceptable.

There were no studies that directly compared the effect of the BLVR procedure on lung volume reduction as a means to examine its safety. However, when considering the severity of the patient and ethical issues, direct comparison was possible. It can be recommended that indirect comparison or long-term observational studies are used to determine safety and effectiveness.

**Conclusion**

Our results suggest that BLVR may be an effective and safe procedure for the treatment of severe COPD patients with emphysema, based on existing studies. Furthermore, the intervention showed lung function improvement in patients with complete fissure, whereas patients with incomplete fissure showed no response. Therefore, more studies are needed to further investigate target patient selection and patient safety.

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**Disclosure**

The authors have no conflicts of interest to disclose.

**References**
