**LUNG VOLUME REDUCTION SURGERY: AN OVERVIEW**

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**INTRODUCTION**

Lung volume reduction surgery (LVRS) continues to be one of the most controversial cardiothoracic procedures in recent years. The report by Cooper et al. in 1995 showing the benefits from lung volume reduction surgery by improving pulmonary function, was rapidly disseminated throughout the United States and the world. It modified the approach of Brantigan and Mueller by using a median sternotomy, thus allowing access to both lungs and used a buttressed staple excision technique. The idea was then to remove the hyperinflated and functionless “target areas” resulting in improvement of the remaining lung.

During early experience with LVRS, functional results, operative mortality as well as morbidity were highly variable. Questions regarding validity of the early clinical reports, incomplete follow-up bias, selection criteria and survival, confounded the interpretation of clinical data on LVRS. Patients with upper, lower and diffuse distribution of emphysema were included; we also analyzed as key points perioperative morbidity and mortality and lung function measurement as FEV1. Bullous emphysema was excluded from this review. Surgical approach included median sternotomy, unilateral or bilateral thoracotomy, and videothoracoscopy with stapled or laser ablation. Results of prospective randomized trials between medical management and LVRS are essential before final assessment can be established.

**METHODS**

We conducted a methodological assessment of the literature, where studies published in English were included. Studies on lung volume reduction surgery were identified using Pubmed (MEDLINE) and Cochrane Library literature in English. Search words such as lung volume reduction surgery or lung reduction surgery, pneumoplasty or reduction pneumoplasty, COPD or chronic obstructive pulmonary disease and surgery, were used. We also compared medical therapy and surgical technique. Studies consisting of randomized controlled trials, controlled clinical trials (randomized and nonrandomized), reviews and case series were analyzed. Questions regarding validity of the early clinical reports, incomplete follow-up bias, selection criteria and survival, confounded the interpretation of clinical data on LVRS. Patients with upper, lower and diffuse distribution of emphysema were included; we also analyzed as key points perioperative morbidity and mortality and lung function measurement as FEV1. Bullous emphysema was excluded from this review. Surgical approach included median sternotomy, unilateral or bilateral thoracotomy, and videothoracoscopy with stapled or laser ablation. Results of prospective randomized trials between medical management and LVRS are essential before final assessment can be established.

**SUMMARY**

This study intends to review the literature on the efficacy, safety and feasibility of lung volume reduction surgery (LVRS) in patients with advanced emphysema. Studies on LVRS from January 1995 to December 2009 were included by using Pubmed (MEDLINE) and Cochrane Library literature in English. Search words such as lung volume reduction surgery or lung reduction surgery, pneumoplasty or reduction pneumoplasty, COPD or chronic obstructive pulmonary disease and surgery, were used. We also compared medical therapy and surgical technique. Studies consisting of randomized controlled trials, controlled clinical trials (randomized and nonrandomized), reviews and case series were analyzed. Questions regarding validity of the early clinical reports, incomplete follow-up bias, selection criteria and survival, confounded the interpretation of clinical data on LVRS. Patients with upper, lower and diffuse distribution of emphysema were included; we also analyzed as key points perioperative morbidity and mortality and lung function measurement as FEV1. Bullous emphysema was excluded from this review. Surgical approach included median sternotomy, unilateral or bilateral thoracotomy, and videothoracoscopy with stapled or laser ablation. Results of prospective randomized trials between medical management and LVRS are essential before final assessment can be established.

**KEY WORDS:** Thoracic surgery. Thoracic surgery, video-assisted. Pulmonary emphysema. Pulmonary disease, Chronic obstructive. Video-assisted surgery.

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In order to be included in this review, studies consisted of randomized controlled trials, controlled clinical trials (randomized and nonrandomized), reviews and case series. Entry criteria for studies included: perioperative morbidity and mortality; lung function measurement as FEV1; patients with any kind of heterogeneous emphysema (upper, lower or diffuse) while; bullous emphysema was excluded from this review. The surgical approach included median sternotomy, unilateral or bilateral thoracotomy, and videothoracoscopy with stapled or laser ablation. The authors attempted to view LVRS as a surgical option for patients with severe emphysema - with particular emphasis placed on safety and efficacy aspects of the procedures - also offering a critical review of this technique.

**Physiopathology of Emphysema - Understanding the Problem**

This disease, which is part of a spectrum of conditions also known as chronic obstructive pulmonary disease (COPD), is frequent in smoker patients, usually in association with features of chronic bronchitis. Its physiopathology is mainly characterized by airflow obstruction and hyperinflation. Briefly, it is defined anatomically by “abnormal, permanent enlargement of airspaces distal to the terminal bronchiole, accompanied by destruction of their walls and without obvious fibrosis.”

Remodeling of the peripheral lung units produces a decreased elastic recoil pressure. As a result, under static conditions, the emphysematous lung requires less pressure to inflate, but once inflated it exerts less emptying pressure than a normal lung. There is as well, a significant mechanical compromise of the diaphragm (flattened position in chest radiographs), resulting in a markedly increased work of breathing.

The diaphragm dysfunction is a main problem while the disease worsens: affecting chest wall mechanics, resulting in further increase of the work of breathing. Due to its flattened position, a result of over stretch of its fibers, the diaphragm in less capable of generating inspiratory force. As such, the less effective diaphragm work results in fatigue and respiratory failure.

Airway inflammation, bronchospasm and increased secretion also have a role in the increased airway pressure. The clinical consequence is incomplete exhalation with intrinsic positive-end-expiratory pressure (auto-PEEP) or dynamic pulmonary hyperinflation caused by retained gas volume in the lungs.

**Medical and Surgical Therapies**

**Medical Treatment**

The main goal of medical therapy for emphysema patients are to: retard chronic disease progression; treat acute exacerbations; control symptoms and improve quality of life. Medical management includes patient education and risk factor avoidance, by eliminating causative agents (tobacco), prevention of infection, pulmonary toilet, rehabilitation and pharmacologic treatment.

Current management of COPD can be divided into pharmacologic and nonpharmacologic categories, according to the Global Initiative for Chronic Obstructive Lung Disease 2003 (GOLD 2003). Pharmacologic treatment include bronchodilators, inhaled corticosteroids, combination therapy and long-term oxygen therapy. Non-pharmacologic therapies include smoking cessation, optimizing nutrition, pulmonary rehabilitation, mechanical ventilation and lung volume reduction surgery (LVRS) for selected cases. Current guidelines recommended inhaled long-acting bronchodilators as the main method of therapy. Calverly et al. performed bronchodilator combination testing in 660 COPD patients classified according to the European Respiratory Society (ERS) and the American Thoracic Society (ATS) spirometric parameters and showed that 55% of patients changed from irreversible to reversible status. Therefore, reduction or elimination of dependence on systemic corticosteroids should be an essential goal of a rehabilitation program. Some authors postulated that randomized trials have failed to show a significant effect of inhaled corticosteroids on pulmonary function, but a meta-analysis by Sutherland showed that high-dose inhaled corticosteroids reduced decline of FEV1, when compared with placebo.

Cigarette smoking cessation should be the physician’s first intervention, especially if there is some indication for surgical treatment. It is the only long-term intervention that shows clear evidence in lung function improvement furthermore it is a demonstration of the patient’s commitment to treatment. Supplemental oxygen therapy is the only approach that increases lung function and survival in COPD patients, as shown in randomized trials. Patients with a PaO2 < 55 mmHg or a SaO2 < 88% should receive supplemental oxygen therapy; if the patient has a PaO2 55-59 mmHg or a SaO2 < 89% with signs of pulmonary hypertension or cor pulmonale, supplemental oxygen is also indicated.

Exercise training or pulmonary rehabilitation aim to optimize performance of daily living activities and maximal exercise tolerance. Mechanisms of lung functional improvement and exercise tolerance have not been fully established, but effects such as improved motivation, improved muscle function and biomechanics, desensitization to dyspnea and increased aerobic capacity are evident.

**Surgical Treatment**

Because medical therapy has been relatively ineffective in slowing the progression of emphysema, surgical attempts have occurred in the past, both to palliate and treat this disease. Early maneuvers included costochondrectomy and transverse sternotomy to improve thoracic mobility; other authors attempted to limit lung expansion with thoracoplasty and phrenectomy or elevate the diaphragm with pneumoperitoneum or abdominal constrictive belts.

In the 1950’s Otto Brantigan and Mueller speculated about the wedge excision of emphysematous patients thus decreasing the lung volume, leading patients to reduce dyspnea. This idea was based upon the concept that a smaller lung could restore the efficiency of the respiratory pump. Indeed, Brantigan’s procedure did produce some clinical improvements in pulmonary function, but the mortality rate was high (18%) and the procedure was soon abandoned. Thirty years later, Cooper and cols., revitalized this procedure with an operation known as lung volume reduction surgery or LVRS. This operation involved resection of 20% to 30% of hyperinflated lung that should be noncontributory to effective ventilation. The most affected portions were excised using a linear stapling device fitted with strips of bovine pericardium to buttress the staple lines and eliminate air leakage through the stapled holes. Expected benefits were improvement in elastic recoil, lung compliance and chest wall conformation to reduce hyperinflation and allow resumption of a more normal diaphragmatic position. Improvement in ventilation-perfusion matching in the remaining lung tissue leading to better pulmonary function should occur as well. Their first report showed no early or late mortality related to the procedure.
Initial studies reported significantly positive clinical results, which led LVR to be performed by many surgeons worldwide. Short-term results from many nonrandomized studies have been reported. Most of them showed reduced hyperinflation and improved pulmonary function and also better ventilatory mechanics and exercise tolerance at 3 and 6 months after surgery. Because LVR is a palliative, elective procedure, one of the major concerns was the related mortality rate. The selection process included marked hyperinflation of the chest and sufficient variation in the emphysema, to provide target areas accessible to lung resection. The degree of regional parenchymal destruction is better analyzed by computed tomography of the chest and distribution of function should be assessed by radionuclide ventilation-perfusion lung scanning. The latter procedure, our subject of interest, represents 20-30% of volume reduction in one or both lungs by means of stapler resection, laser application, or both. Acceptable results from stapler resection have been reported through a sternotomy, thoracotomy, clam shell incision and thoracoscopy. There have been some controversial results regarding the type of operation and whether this should be applied unilaterally or bilaterally. The most extensive comparison of LVRs by median sternotomy (MS) or video assisted thoracic surgery (VATS) was published by the National Emphysema Treatment Trial (NETT). They analyzed 343 patients in the MS arm and 146 patients in the VATS arm and found that the two approaches carry similar risks of 30-day, 90-day and overall mortality. Although there was a slight trend for higher mortality after MS than VATS, this was not statistically significant. Both approaches determine similar changes in exercise capacity, lung function and disease specific quality of life, also showing significant improvement after 6 months. The mean hospital and physician costs for the LVRs admission were $8,207 less for the VATS group compared to the MS group (95% confidence interval[CI]; $0–0,03). Mean total costs during the 6 months following surgery were $10,428 lower for the VATS group (95% CI on difference; $0–0,005). Costs analysis reflects fewer Intensive Care Unit stays and a reduced overall length of stay for the VATS group. This important report concluded that choice of the approach is a matter of the surgeon’s preference and experience.

Functional status before and after surgery is assessed by measuring multiple parameters of pulmonary function and quality of life indicators, but FEV1 is the most used as single indicator of functional status. Using this method, stapling is usually associated with more short-term (3-6 months) improvement than the laser technique. With regard to the staple line and comparisons of the type of buttressing (bovine pericardium or collagen), no significant differences have been found in the efficacy. Another controversial question is if LVRs should be attempted unilaterally or bilaterally. Unilateral LVR should be performed intentionally in patients with distinct heterogeneity of emphysema.

between lungs identified and graded by radiologic findings, whereas simultaneous bilateral LVR is preferred for patients with heterogeneous disease in both lungs but symmetrically distributed between the lungs. Spirometry, lung volumes and quality of life appeared to be superior for bilateral compared to unilateral LVR, although there was no significant difference in mortality between the two methods. Sema and colleagues found that survival at 2 years was better for unilateral than unilateral VATS LVR. A recent study published by Pompeo and colleagues analyzed 97 patients with upperlobe prevailing emphysema by unilateral LVR, and showed 82% of 5-year related survival.

There is an important question regarding the surgical technique: it has changed from an inverted U-shaped that goes from an anterior aspect of the upper lobe toward the apex and then down the back. The current technique is just a resection from the front, straight toward the back, removing almost all the upper lobe on the right. On the left, the portion of the upper lobe is removed almost completely, thus just the lingula is left intact. Although this transverse resection is now preferred over oblique resection and it seems that there is no significant difference between both methods. This transition has been gradual, therefore there is no way to identify patients retrospectively with regard to the type of stapled line.

Irrespective of the method of choice, this operation is to be considered after maximal medical therapy has failed to produce satisfactory palliation and intended to return the emphysematous lung to an earlier stage in the natural history of obstructive lung disease.

**DISCUSSION**

The American Thoracic Society has classified LVR as an innovative rather than experimental procedure, however such a technique is surrounded by many unresolved questions. Until these questions are answered, LVR cannot be considered a standard therapy. One of the major issues in LVRs is how to select appropriate candidates and how to assess and interpret results.

The current position of the American Thoracic Society is that LVRs should be performed only at centers where it can be completely studied through clinical trials and extensive physiologic evaluations. Appropriate candidates are those with severe emphysema refractory to medical therapy, disabling symptoms and evidence of air trapping (FEV1 < 35%). Although patients who have FEV1 > 40% are probably not sufficiently incapacitated for surgery, there is no consensus on a lower limit of FEV1, as an exclusion criterion for LVRs. When the FEV1 is less than 30% of the predicted, 50% of patients will die within 3 to 4 years.

Many centers have patients over 75 years of age and significant associated illnesses, such as coronary artery disease or morbid obesity. Preoperative evaluation should comprise complete pulmonary and cardiac tests and other tests, such as dynamic MRIs, positron emission tomography scan and sleep studies could be used for screening. The distribution of emphysema is classified as heterogeneous or homogeneous based on the high-resolution computed tomography under a visual score system. The radiologist classifies the craniocaudal distribution of emphysema as predominately affecting the upper lobes, predominantly affecting the lower lobes, diffuse or predominantly affecting the superior segments of the lower lobes; the latter three categories were classified as non-upper lobe emphysema. Before randomization eligible patients completed 6 to 10 weeks of pulmonary rehabilitation supervised by the transplant.
team. According to the NETT, patients with a FEV$_1$ 20% or less and an either non-upper-lobe emphysema or a carbon monoxide diffusing capacity that was 20% or less of the predicted value, were determined to be at high risk of death after LVRS, with a low probability of functional benefit.

The National Emphysema Treatment Trial (NETT) was designed and supported by the National Heart and Medical Organization (Medicare and Medicaid Services) since January 1996, when Medicare stopped funding for this procedure. The NETT growth as a multicenter, randomized and large scale clinical trial to evaluate the effects of LVRS and to determine those who would and who would not benefit from this procedure. The main purposes were to compare medical to surgical therapy with respect to short and long term improvement in lung function and quality of life, to determine whether different surgical approaches (median sternotomy or VATS) are related to different outcomes and finally to evaluate costs associated to this procedure. Similar studies are being conducted in Massachusetts, Canada, and Great Britain.

The first NETT results examined 1033 patients and identified a group of 69 patients at a high risk of death compared with medical management in a group of patients with a low preoperative FEV$_1$ and a uniform pattern of emphysema or low D$_{LCO}$. This report showed improved outcomes related to emphysema heterogeneity, upper versus lower lobe LVRS and bilateral versus unilateral LVRS.

In 2003, NETT reported the effects of LVRS versus medical therapy on survival and maximum exercise capacity in 1218 patients who were randomized for treatment between January 1998 and July 2002 and monitored for a mean of 2.4 years (figure 1). A subgroup of high risk patients (n = 140) - homogeneous pattern of emphysema on chest CT, VEF$_1$ $\leq$ 20% and D$_{LCO}$ $\leq$ 20% - also was analyzed (figure 2). The 90-day mortality rate in the surgery group was 7.9%, higher than the medical therapy group ($p < 0.001$). Additional outcomes reported included pulmonary function, oxygen requirement, distance walked in 6 minutes, quality of life and respiratory symptoms. These data were updated in 2006, with a mean follow-up of 4.3 years.

The NETT studies concluded that lung volume reduction surgery improves exercise capacity, but does not confer a survival advantage over medical therapy. It does provide a survival advantage for patients with both predominantly upper-lobe emphysema and low baseline exercise capacity.

Meyers$^{25}$ and cols. published results from a group of 20 patients with a FEV$_1$, and diffusion capacity of carbon monoxide of 20% or less who underwent bilateral LVRS and showed a 90-day operative mortality of 5%. In all patients the FEV$_1$ increased from 0.46 (17%) to 0.78 (32%), a 73% change; the D$_{LCO}$ increased from 16% to 27%, a 70% improvement and room air PaO$_2$ increased from 55 mmHg to 64 mmHg. The Kaplan-Meier 5-year survival rates did not differ between the high-risk and non-high-risk patients. He concluded that patients in this selected group might experience improvements in lung function, exercise tolerance and quality of life with acceptable morbidity and mortality after LVRS. Paradoxically this group achieved an improvement greater than in the rest of series, possibly explained by the larger target areas and lower FEV$_1$.

It is indeed well established that these high-risk patients represent a relative contraindication to such procedure. The NETT$^{1}$ reported two high risk groups randomized for surgical treatment: patients with low FEV$_1$, and homogeneous distribution of emphysema and patients with very low FEV$_1$, and a low D$_{LCO}$. The overall mortality in this group was 28.6% compared to 7.9% in the non-high-risk patients. These findings corroborate observations from early LVRS experience, and have added additional weight of evidence to the belief that these patients (homogeneous pattern of emphysematous destruction) are poor candidates. Also interesting in the NETT study, is the fact that the exercise capacity after 24 months had improved in 16% of patients in the surgery group as compared to 3% of patients in the medical group ($p < 0.001$). Among the 610 patients assigned to medical therapy, the 90-day mortality rate was 1.9%, 33 (5.4%) underwent LVRS outside the study and 15 (2.5%) received lung transplantation during follow-up, may be related to worsening of lung function or lack of alternative therapy available. Miller and cols.$^{26}$ published a recent study comparing LVRS to optimal medical therapy in two clinical trials and concluded that in carefully selected patients with advanced heterogeneous emphysema, quality of life had improved 6 to 12 months after LVRS combined with optimal medical therapy, results superior to those of patients who received optimal medical therapy only. Fishman and cols.$^{2}$ concluded that patients with upper-lobe predominant emphysema and low exercise capacity had improved survival with LVRS, when compared to the medical therapy group and indicated two outcome predictors: distribution of emphysema and exercise capacity following pulmonary rehabilitation. Pulmonary rehabilitation provides substantial benefits for patients with severe COPD, including increased exercise capacity and decreased dyspnea. Although many patients never participate in this therapy modality, it is certainly less costly than surgery and associated with less immediate morbidity and mortality$^{27}$.

Use of LVRS with undiagnosed pulmonary nodules and/or lung cancer is controversial and a wide range of abnormal tissue including adenocarcinoma, bronchoalveolar and carcinoids tumorlets has been found in lung resection following LVRS. De Rose$^{29}$ and cols. advocate LVRS combined with nodule resection and reported recurrent lung cancer in 1 of 14 patients one year after operation by this technique. LVRS also has been performed with coronary artery bypass surgery, cardiac valvular and aortic aneurysm surgery.

Another controversial point is the mortality rate reported in some series; a few authors published hospital mortality, others operative mortality and others 30-day related mortality. Because significant mortality occurs between 30 and 90 days, it is established that 90-day is the most useful indicator. A recent meta-analysis published by Berger$^{30}$ and colleagues showed similar 6 and 12 months mortality between LVRS and medical groups, after random assignment to treatment. It is well known that patients with predominantly upper-lobe have lower mortality and greater chance of improvement in exercise capacity than patients with non-upper lobe emphysema$^{30,31}$.

LVRS has been employed adjunctively in these other special conditions:

1. resection for stage I lung cancer;
2. to wean ventilator dependent COPD patients;
3. to reduce unilateral hyperinflation of the native lung after single lung transplantation;
4. to serve not only as a bridge procedure but as an alternative for patients awaiting lung transplantation$^{31}$.

**Conclusion**

The conclusion of this review is based upon the important benefits that LVRS can provide to patients under careful selection and rigorous preoperative and perioperative care. How long these benefits will last...
O objetivo deste estudo é revisar a literatura acerca da eficácia, segurança e viabilidade da cirurgia reductora de volume pulmonar (CRVP) em pacientes com enfisema pulmonar avançado. Estudos de CRVP de janeiro de 1995 a dezembro de 2009 foram incluídos através de pesquisa na PubMed (MEDLINE) e Cochirane Library, na literatura inglesa. Palavras de busca tais como lung volume reduction surgery (LVRS) ou lung volume reduction therapy foram utilizadas. Também realizamos comparação entre terapia médica e cirúrgica. Os estudos analisados consistiam de randomizados controlados, estudos clínicos controlados, (randomizados e não randomizados), revisões e séries de casos. As questões acerca da validade através dos relatos iniciais, seguimentos incompletos, critérios de seleção indefinidos e análises de sobrevida confundiram a interpretação dos dados clínicos provenientes da CRVP. Pacientes com enfisema de predominio em lobos superiores, inferiores e difuso, foram incluídos; também analisamos pontos chave, tais como morbidade e mortalidade peri-operatórias, assim como a medida da função pulmonar através do VEF 1. Enfisema tipo bolhoso foi excluído desta revisão. Foram incluídas para análise também vias de acesso cirúrgico como esternotomia mediana, toracotomias unilateral ou bilateral e videotoracoscopia unilateral ou bilateral com grameamento ou ablação por laser. Os resultados dos estudos prospectivos randomizados entre o tratamento clínico e a CRVP são essenciais para que alguma conclusão possa ser definitiva. (Rev Assoc Med Bras 2010; 56(6): 719-23)

Relações de interesse: Nenhuma

Referências