Practice Parameters for the Use of Laser-Assisted Uvulopalatoplasty: An Update for 2000

Michael Littner MD,¹ Clete A. Kushida MD, PhD,² Kristyna Hartse PhD,³ W. McDowell Anderson MD,⁴ David Davila MD,⁵ Stephen F. Johnson MD,⁶ Merrill S. Wise MD,⁷ Maxwell Hirshkowitz PhD,⁸ and B. Tucker Woodson MD, FACS⁹

¹VA Greater Los Angeles Healthcare System, and UCLA School of Medicine, Sepulveda, CA; ²Stanford University Center of Excellence for Sleep Disorders, Stanford, CA; ³Sleep Consultants, Inc., Fort Worth, TX; ⁴College of Medicine, University of South Florida, Tampa, FL; ⁵Baptist Medical Center, Little Rock, AR; ⁶Saint Patrick Hospital Sleep Center, Missoula, MT; ⁸Departments of Pediatrics and Neurology, Baylor College of Medicine, Houston, TX; ⁸Baylor College of Medicine, Houston VAMC Sleep Disorders and Research, Houston, TX; ⁹Department of Otolaryngology and Communication Sciences, Medical College of Wisconsin, Milwaukee, WI

Summary: Laser-assisted uvulopalatoplasty (LAUP) is an outpatient surgical procedure which is in use as a treatment for snoring. LAUP also has been used as a treatment for sleep-related breathing disorders, including obstructive sleep apnea. The Standards of Practice Committee of the American Academy of Sleep Medicine reviewed the available literature, and developed these practice parameters as a guide to the appropriate use of this surgery. Adequate controlled studies on the LAUP procedure for sleep-related breathing disorders were not found in peer-reviewed journals. This is consistent with findings in the original practice parameters on LAUP published in 1994. The following recommendations are based

on the review of the literature: LAUP is not recommended for treatment of sleep-related breathing disorders. However, it does appear to be comparable to uvulopalatopharyngoplasty (UPPP) for treatment of snoring. Individuals who are candidates for LAUP as a treatment for snoring should undergo a polysomnographic or cardiorespiratory evaluation for sleep-related breathing disorders prior to LAUP and periodic postoperative evaluations for the development of same. Patients should be informed of the best available information of the risks, benefits, and complications of the procedure.

INTRODUCTION

LASER-ASSISTED UVULOPALATOPLASTY (LAUP) HAS BEEN PROMOTED AS A TREATMENT OF SNORING, AND IN SOME CASES, for sleep-related breathing disorders including obstructive sleep apnea (OSA). This surgical procedure is typically performed in an outpatient setting with local anesthesia and without postoperative hospitalization. At the present time, LAUP is in current use. In this article, we review the appropriate patient evaluation and the effectiveness, potential risks, and complications of LAUP for OSA, and provide recommendations for its use. This update generally examines evidence for LAUP in the therapy of OSA since the publication of the expert review; grades the evidence available; and modifies and replaces the 1994 practice parameters.

METHODS

Medline searches for articles on LAUP were conducted through September 2000. Key words for the search included LAUP, laser-assisted uvulopalatoplasty, laser-assisted uvuloplasty, laser surgery, somnoplasty, base of the tongue reduction, uvulopalatopharyngoplasty (UPPP), uvulopalatoplasty, uvuloplasty, uvulotomy, uvula, and all possible combinations of the preceding terms with snoring, obstructive sleep apnea, sleep apnea syndromes, and upper airway surgery. This search led to a total of 641 articles. Thirty-two of these articles were published prior to the original American Academy of Sleep Medicine's

Accepted for publication April 2001

Address correspondence to: Standards of Practice Committee, American Academy of Sleep Medicine, 6301 Bandel Road, Suite 101, Rochester, MN 55901; Tel: 507-287-6006; Fax: 507-287-6008; E-mail: aasm@aasmnet.org

(AASM) Practice Parameters for the Use of Laser-Assisted Uvulopalatoplasty¹ in 1994, which incorporated 17 of the 32 articles in that previous review of the literature. Articles in all languages were considered for inclusion, and were screened based on their English-language abstracts. A total of 123 articles were identified as potentially relevant based on review of the abstracts. Of these, 90 were obtained in full length and examined. Upon review of these articles, an additional 45 references were discovered by pearling (i.e., the process of selecting relevant articles referenced in the original article). These were references located in publications not typically found through Medline. The types of these publications, with the total number of publications per type (in parentheses) are listed: books (6), coursebooks (1), meeting and symposium abstracts or proceedings (8), highly specific or trade journals (30). Articles entered into the evidence tables (Tables 1 and 2) included randomized trials and nonrandomized controlled or concurrent cohort studies on the comparison with UPPP for snoring and OSA (Table 1) and peer-reviewed case series and historical cohort studies on the efficacy of LAUP for OSA (Table 2), with a minimum of five patients and a clearly defined outcome that could be used to adequately assess the therapy. In the case of the peer-reviewed case series and historical cohort studies entered in Table 2, studies were included only if the "effect size" (Table 3) or the overall effect of LAUP on the number of respiratory events during sleep (described below) could be derived from the article. Articles describing nonrandomized historical cohort studies (13), case series (45), and other studies (69) derived from the search were found useful as background articles. The Standards of Practice Committee's levels of evidence (Table 4) for treatment-related evidentiary articles, which are used to support the strength of the recommendations (Table 5) in this paper, are found in the evidence tables (Tables 1 and 2).

On the basis of this review and noted references, the Standards of Practice Committee of the American Academy of Sleep Medicine, in conjunction with specialists and other interested parties, developed the review and recommendations included in this paper. In most cases, the conclusions are based on evidence from studies published in peer-reviewed journals that were evaluated as noted in the evidence tables (Tables 1 and 2). However, when scientific data are absent, insufficient, or inconclusive, the recommendations are based upon consensus opinion. The strength of each recommendation is based on the level of the evidence available or on consensus when evidence is lacking.

The Board of Directors of the American Academy of Sleep Medicine approved this review and these recommendations. All authors of this review, members of Standards of Practice Committee, and the Board of Directors completed detailed conflict-of-interest statements and were found to have none with regard to this subject.

These practice parameters define principles of practice that should meet the needs of most patients in most situations. These guidelines should not, however, be considered inclusive of all proper methods of care or exclusive of other methods of care reasonably directed toward obtaining the same results. The ultimate judgment regarding the propriety of any specific care must be made by the physician in light of the individual circumstances presented by the patient and the available diagnostic and treatment options as resources.

The American Academy of Sleep Medicine expects these guidelines to have a positive impact on professional behavior, patient outcomes and, possibly health care costs. These practice parameters reflect the state of knowledge at the time of development and will be reviewed, updated, and revised, as new information becomes available.

Background

LAUP is a surgical procedure that typically relies on the use of a carbon dioxide (CO₂) laser to vaporize the uvula and a part of the free edge of the soft palate during one to several sessions. Within the scope of this definition, various degrees of tissue are ablated using slightly different techniques. This procedure is different from conventional uvulopalatopharyngoplasty (UPPP), in that LAUP is performed during a comparatively brief surgical session, reduces far less palatal tissue and does not alter the tonsils or the pharyngeal pillars, uses a laser rather than a scalpel, requires no wound closure, uses local rather than general anesthesia, is conducted in an ambulatory rather than hospital setting, and requires no postoperative hospital stay.1 LAUP is distinguished from the laser palatoplasty procedure described by Ellis² in which a soft palate lesion produced by a neodymium:yttriumaluminum-garnet laser induces scarring, which stiffens the soft palate and reduces "palatal flutter," which in turn, reduces snoring.

Patient Evaluation

The selection process for candidates for this procedure ranges from patient history, questionnaire data, use of the Müller maneuver, oral and nasopharyngoscopic examination, polysomnography, and a variety of imaging studies. Although some investigators proposed decision algorithms³ or imaging studies to localize

the site of obstruction,^{4,5} there is no consensus on the preoperative selection process for this procedure. However, a patient deciding on LAUP as a treatment for snoring should be properly screened for a more severe sleep-related breathing disorder such as OSA. Clinical evaluation can be unreliable; a clinical history and results of a physical examination by a physician to generate a subjective judgment as to whether a given patient did or did not have OSA yielded a correct identification in 52% of patients with OSA and a specificity of 70%.6 Another study showed that out of 73 patients seeking LAUP treatment, 69 (95%) had OSA by polysomnography, even though 41% presented only with a complaint of snoring.⁷ Additionally, the patients' subjective ratings of snoring loudness, frequency, and consequences did not correlate with any of the respiratory variables obtained by polysomnography. Thus, a sleep study, in the form of standard polysomnography or Level III recording also called a cardiorespiratory study,8-¹⁰ is indicated to exclude the possibility of OSA in potential candidates for this procedure for snoring. A Level III recording includes at least four channels with recording of at least two respiratory effort channels or a respiratory effort channel and an airflow channel, plus oximetry and either heart rate or electrocardiogram.

Effectiveness, Risks, and Complications of LAUP for Snoring and OSA

In 1990, Kamami described the use of LAUP on 31 adult patients.¹¹ Following up to seven sessions a maximum of three weeks apart, snoring was completely eliminated or remained as an occasional soft snore in 24/31 (77.4%) of the patients, and a persistent non-disturbing snore in 7/31 (22.6%) of the patients. Neither infection nor significant bleeding was detected; patients reported pain similar to a simple "sore throat." The patients noted improvement in fatigue, morning headaches, and irritability; however, it is unknown whether any of the subjects had OSA, since preoperative screening polysomnography was not performed.

Although there are a number of case series subsequent to Kamami's original study, randomized placebo-controlled studies on the effectiveness of LAUP for OSA are lacking. This lack provides evidence of limited value in determining if LAUP has efficacy in OSA. However, by combining a number of the case series studies, 5,12-17 it is possible to determine an overall effect of LAUP on the number of respiratory events during sleep. The "effect size" of each study is derived from the difference between the pre- and post-LAUP number of apneas and hypopneas per hour of sleep (also called the pre- and post- apnea hypopnea index, AHI) divided by the standard deviation of the pre-LAUP AHI.18 The effect size can be adjusted by a factor related to the number of subjects in each study.¹⁹ The overall effect of a number of studies can be expressed as the average of the sum of individual unadjusted or adjusted effect sizes^{18,19} of each study. The case series studies were selected from the total number of case series articles obtained through our literature search. The criteria used for inclusion of these articles in the calculation of effect size were studies in which the mean pre- and post-LAUP AHI across subjects as well as the pre-LAUP standard deviation were provided in the article, or could be derived from data present in the article. When the effect size analysis was performed (Table 3), the average unadjusted effect size was 0.392. The average adjusted effect size was 0.251. Because there is no comparison with placebo or with another procedure, it is difficult to determine if this effect is likely to be meaningful. However, in general, an effect size between 0.2 and 0.5 (as is the average in the LAUP studies) is considered to be in the small range. By comparison, an effect size between 0.5 and 0.8 is considered to be medium and greater than 0.8 is considered to be large.

The reader should be aware of the following in interpreting information on LAUP. It is not clear if the general interpretation of effect size can be applied to the specific case of LAUP. The reduction in AHI may not be clinically significant since there are few outcome measures such as sleepiness and systematic quality of life reported in the literature. Although the overall effect is a small improvement, individual patients may show no reduction or an increase in AHI.²⁰ Apart from the near-term post-operative effects of LAUP on AHI, the long-term efficacy of LAUP on OSA is undefined. Interpretation of the effect of LAUP is based on studies that have described different surgical procedures ranging from excising comparable amounts of tissue as those removed with UPPP,²¹ to varied and lesser excisions.^{22,23}

As illustrated in Table 2, there are six Level III studies, representing nonrandomized controlled or concurrent cohort studies, 3,21,24-27 comparing LAUP vs. UPPP (either with or without tonsillectomy).3,21,24-27 One study evaluated OSA,24 one study examined snoring and OSA3 and one study examined snoring and upper airway size.²⁶ Two of the three studies showed a decrease in AHI which because of sample size could not be compared for degree of efficacy to UPPP;3,24 the remaining study showed worsened postoperative upper airway anatomic characteristics by oral and nasopharyngoscopic examination for LAUP compared to UPPP patients.²⁵ Four studies reported subjective postoperative improvement in snoring levels with LAUP and no significant differences in levels of improvement between LAUP vs. UPPP.^{3,21,26-27} However, interpretation of the results of all of the above studies is difficult given the relative lack of detailed statistical analyses of the data. As mentioned above, comparisons between studies are further limited by lack of standardization of the procedure.

Lastly, the long-term effectiveness of LAUP on treatment of snoring has not been convincingly established. Two separate studies found snoring improvement of 89.6% and 90%, in patients assessed between one and eight years and at five years following LAUP.28,29 Less satisfactory results were found in a study that showed snoring improvement was reduced to 62.2% beyond two years.¹⁴ Another study found that 22% of patients had recurrence of snoring between 18 and 24 months following LAUP, with an overall success rate of 55% at 24 months, 30 and a separate study found snoring improvement in 43% of patients, with 21% showing no improvement and 36% showed significant deterioration on sleep studies performed 3 to 24 (mean=7) months postoperatively.³¹ Following an average post-LAUP duration of four years, another study found that 51.6% of patients reported that their snoring was eliminated. 13 As mentioned, the long-term efficacy on LAUP on OSA is not defined but should be considered problematic in view of the inconsistent findings on the long-term efficacy of LAUP on snoring.

There are data to suggest that the pain levels associated with LAUP may be comparable to those of UPPP. One study showed no difference between the average pain scores for the first (typically the most painful) LAUP stage and UPPP.²⁶ However, the patients treated with UPPP remained in the hospital overnight and received parenteral analgesia. Another study showed similar maximum pain peaks and intensity for LAUP vs. UPPP, with comparable mean durations of the pain period of 13.76 and 11.80 days, respectively.³ Similar results were reported in a separate study, which found comparable mean durations of the pain period for LAUP (13.8 days) vs. UPPP (14.3 days).³²

Besides pain, the most commonly reported side effects from LAUP appear to be transient velopharyngeal insufficiency, minor bleeding, local infection, globus sensation, and minor dysphonia and dysphagia.^{33,34} Based on the literature review, the most common side effects with their reported frequency of occurrence are listed in Table 6. In 27% of LAUP patients, either persistent dysphagia³⁵ or mild or moderate scar fibrosis²⁴ have been observed. Postoperative swelling may compromise an already marginal upper airway; use of narcotics or sedatives may further complicate this problem. Alcohol should be avoided because of its adverse effects on upper airway muscle tone and closing pressures in snorers.³⁶ The smoke plume from lasers can create a biological and chemical hazard for the patient and surgical team; however, an efficient smoke evacuator used during LAUP can obviate this hazard.³⁷

There is also evidence to indicate that LAUP may result in a diminished velopharyngeal air space and decreased distensibility.²⁵ This study suggests that these structural modifications of the upper airway may decrease airway resistance, resulting in further narrowing during inspiration and collapse of the upper airway at the level of the tongue base, and consequent OSA. These results, from an anatomical perspective, indicate that LAUP may have a worse outcome than UPPP. A separate study examining LAUP patients between 48 and 72 hours after LAUP found worsening of the AHI, with a significant decrement in the cross-sectional area of the airway by videoendoscopy.²⁰ A study examining histopathologic changes of the soft palate after LAUP found extensive thermal-induced changes including diffuse fibrosis, oral epithelia ulceration, and a patchy inflammatory reaction, which the authors speculate may be responsible for worsening of OSA.38

The selection process for candidates for LAUP or the anatomic, histopathologic, and physiologic effects of this procedure have not been well characterized, and there is a lack of understanding of its consequences on pathologic respiration and its long-term effectiveness. In general, since insufficient data exists on the effectiveness and risks of LAUP, patients who elect to undergo this procedure as a treatment for snoring should have appropriate preoperative evaluation including screening for OSA, and should have close postoperative follow-up to monitor the patient for possible complications of this procedure.

CONCLUSIONS AND RECOMMENDATIONS

The following recommendations of the Standards of Practice Committee and the Board of Directors of the American Academy of Sleep Medicine are similar to those published in its last report in 1994, since adequate controlled studies on the LAUP procedure were not found in peer-reviewed journals. The classification of evidence was adapted from the suggestions of Sackett³⁹ (Table 4). Recommendations are given as standards, guidelines, and options, as defined in Table 5.

1. LAUP is not recommended for the treatment of the sleep-related breathing disorders including obstructive sleep apnea. (Guideline)

There is insufficient evidence to recommend LAUP for the treatment of the obstructive sleep apnea syndrome. The Level V, Grade C evidence from seven articles^{5,12-17} indicates that LAUP provides a small overall decrease in AHI in a group of patients, that preoperative prediction strategies for selecting patients who respond have not been developed, that some patients may have an increase in AHI, and that there is insufficient information on other outcome measures or long-term efficacy. Therefore, we do not recommend LAUP for the treatment of obstructive sleep apnea. This recommendation is similar to a recommendation of the previous practice parameter paper.¹

2. LAUP is not recommended as a substitute for UPPP in the treatment of sleep-related breathing disorders including obstructive sleep apnea. (Guideline)

There are three studies with Level III, Grade C evidence^{3,24,25} on comparison including measurement of AHI or airway size. When considered in conjunction with the small effect size of LAUP on AHI, these studies provide insufficient evidence to indicate that LAUP is an acceptable substitute for UPPP with respect to either effectiveness or side effect profiles as a treatment for OSA. This is a new recommendation.

3. LAUP appears comparable to UPPP in relieving subjective snoring. (Guideline)

There are 4 Level III, Grade C studies that compare LAUP to UPPP for snoring. These studies suggest that LAUP can reduce snoring measured by subjective criteria to a similar degree as UPPP. This is a new recommendation.

4. Surgical candidates for LAUP as a treatment for snoring should undergo a preoperative clinical evaluation and a polysomnographic or a cardiorespiratory study⁸⁻¹⁰ to determine if the candidate has a sleep-disordered breathing disorder including obstructive sleep apnea. (Standard)

Since snoring is a primary diagnostic symptom, patients who undergo LAUP should be informed of the need for periodic evaluation for subsequent development of obstructive sleep apnea even if the procedure reduces or eliminates snoring. (Standard)

These recommendations are based on information regarding the natural course of OSA. Snoring may predate onset of OSA, as well as other symptoms of OSA such as excessive daytime sleepiness.⁴⁰ Although snoring is neither necessary nor sufficient for the diagnosis of a sleep-related breathing disorder, it is frequently an

associated symptom. It is estimated that the occurrence of obstructive sleep apnea ranges from 25% to as high as 95% in snorers.^{8,9} In one study reviewing patients seeking LAUP treatment specifically for snoring, 95% had OSA by polysomnography.⁵ The presence of other risk factors for sleep apnea such as obesity and age, as well as other associated symptoms such as daytime sleepiness and witnessed breathing pauses, increase the risk for concomitant sleep apnea. Given the life-threatening conditions (e.g., myocardial infarction, cardiac failure, stroke) associated with sleep-related breathing disorders and the increased risk for motor-vehicle or industrial accidents secondary to the daytime sleepiness related to sleep-disordered breathing, it is prudent to test for these disorders. Patients who elect to undergo LAUP for the treatment of snoring may also be at risk of incurring a delay in the diagnosis of OSA because snoring may be reduced or eliminated by LAUP. Thus, after LAUP for treatment of snoring, the patient should be notified regarding the possibility of developing OSA, and should be monitored for the occurrence of this disorder. These recommendations are similar to recommendations of the previous practice parameter paper.¹

5. The need for medications that affect respiration during the perioperative period should be assessed during the preoperative clinical evaluation (Standard).

This recommendation is based on consensus of the SPC. the perioperative use of narcotics may pose risks for patients who have undergone LAUP operations; therefore, the need for these medications should be carefully assessed during the preoperative clinical evaluation. Careful clinical judgment should be used when prescribing other pain medications, sedatives, sleeping pills and alcohol during the perioperative period. The rationale is that these medications may blunt respiratory drive. This is especially important since postoperative swelling may reduce the caliber of an already narrowed airway. Alternatives, such as oral or topical non-narcotic pain medications during the perioperative periods, should be used whenever possible, and hypnotics and alcohol should be avoided because of their deleterious effects on upper airway tone. This recommendation is similar to a recommendation of the previous practice parameter paper.¹

6. Patients should be informed of the risks and complications of LAUP. (Standard)

There are studies specifically evaluating the risks and complications of LAUP (Table 6). Any patient electing to undergo LAUP for treatment of snoring should be informed of the potential risks and complications of this procedure. This recommendation is based on the documented risks of LAUP and SPC consensus and is similar to a recommendation of the previous practice parameter paper.¹

RECOMMENDATIONS FOR FUTURE RESEARCH

Investigations to identify the best treatment for snoring or OSA should include well-powered, multicenter clinical trials using randomized study designs with an appropriate endpoint or outcome. The use of objective measures for evaluating outcomes and sham or sub-therapeutic controls is encouraged. Future studies should provide LAUP definitions, long-term effectiveness data, cost-benefit analyses, direct comparison between different treatments, and the impact of treatment on quality of life.

REFERENCES

- 1. Standards of Practice Committee of the American Sleep Disorders Association. Practice parameters for the use of laser-assisted uvulopalatoplasty. Sleep 1994;17(8):744-8.
- 2. Ellis PDB, Fowcs-Williams JE, Shneerson JM. Surgical relief of snoring due to palatal flutter. Ann R Coll Surg 1993;75:286-90.
- 3. Remacle M, Betsch C, Lawson B, Jamart J, Eloy P. A new technique for laser-assisted uvulopalatoplasty: decision-tree analysis and results. Laryngoscope 1999;109:763-8.
- 4. Tsushima U, Antila J, Laurikainen E, Svedström E, Polo O, Kormano M. Digital fluoroscopy before and after laser uvulopalatopharyngoplasty in obstructive sleep apnea. Acta Radiologica 1997;38:214-21.
- 5. Skatvedt O, Akre H, Godtlibsen OB. Continuous pressure measurements in the evaluation of patients for laser assisted uvulopalatoplasty. Eur Arch Ororhinolaryngol 1996;253:390-4.
- 6. Viner S, Szalai JP, Hoffstein V. Are history and physical examination a good screening test for sleep apnea? Ann Int Med 1991;115:356-9.
- 7. Khosh M, Keidar A, Zammit GK, Krespi YP. Self-reported symptoms and polysomnographic findings in laser-assisted uvulopalatoplasty candidates. Otolaryngol Head Neck Surg 1994;43 (Abstract).
- 8. Ferber R, Millman R, Coppola M, Fleetham J, Murray CF, Iber C, McCall V, Nino-Murcia G, Pressman M, Sanders M, Strohl K, Votteri B, Williams A. Portable recording in the assessment of obstructive sleep apnea. Sleep 1994;17(4):378-92.
- 9. Standards of Practice Committee of the American Sleep Disorders Association. Practice parameters for the use of portable recording in the assessment of obstructive sleep apnea. Sleep 1994;17(4):372-7.
- 10. Indications for Polysomnography Task Force, American Sleep Disorders Association Standards of Practice Committee. Practice Parameters for the Indications for Polysomnography and Related Procedures. Sleep 1997;20:406-422.
- 11. Kamami Y-V. Laser CO2 for snoring. Preliminary results. Acta Oto-Rhino-Laryngologica Belg 1990;44:451-6.
- 12. Ryan CF, Love LL. Unpredictable results of laser assisted uvulopalatoplasty in the treatment of obstructive sleep apnoea. Thorax 2000;55:399-404.
- 13. Walker RP, Garrity T, Gopalsami C. Early polysomnographic findings and long-term subjective results in sleep apnea patients treated with laser-assisted uvulopalatoplasty. Laryngoscope 1999;109:1438-41.
- 14. Mickelson SA, Ahuja A. Short-term objective and long-term subjective results of laser-assisted uvulopalatoplasty for obstructive sleep apnea. Laryngoscope 1999;109:362-7.
- 15. Walker RP, Grigg-Damberger MM, Gopalsami C. Laser-assisted uvulopalatoplasty for the treatment of mild, moderate, and severe obstructive sleep apnea. Laryngoscope 1999,109:79-85.
- 16. Utley DS, Shin EJ, Clerk AA, Terris DJ. A cost-effective and rational surgical approach to patients with snoring, upper airway resistance syndrome, or obstructive sleep apnea syndrome. Laryngoscope 1997;107:726-34.
- 17. Walker RP, Grigg-Damberger MM, Gopalsami C, Totten MC. Laser-assisted uvulopalatoplasty for snoring and obstructive sleep apnea: results in 170 patients. Laryngoscope 1995;105:938-43.
- 18. Fredric M.Wolf. Meta-Analysis: quantitative Methods for Research

- Synthesis. Sage Publications, 1986.
- 19. Petitti Diana B. Meta-analysis, decision analysis, and cost-effectiveness. Oxford University Press, 1994.
- 20. Terris DJ, Clerk AA, Norbash AM, Troell RJ. Characterization of postoperative edema following laser-assisted uvulopalatoplasty using MRI and polysomnography: implications for the outpatient treatment of obstructive sleep apnea syndrome. Laryngoscope 1996;106:124-8.
- 21. Carenfelt C. Laser uvulopalatoplasty in treatment of habitual snoring. Ann Otol Rhinol Laryngol 1991; 100:451-4.
- 22. Krespi YP, Keidar A, Khosh MM, Pearlman SJ, Zammit G. The efficacy of laser assisted uvulopalatopharyngoplasty in the management of obstructive sleep apnea and upper airway resistance syndrome. Op Tech Otolaryngol Head Neck Surg 1994;5:235-43.
- 23. Dickson RI, Mintz Dr. One-stage laser assisted uvulopalatoplasty. J Otolaryngol 1996;25:155-61.
- 24. Walker RP, Grigg-Damberger MM, Gopalsami C. Uvulopalatopharyngoplasty versus laser-assisted uvulopalatoplasty for the treatment of obstructive sleep apnea. Laryngoscope 1997;107:76-82.
- 25. Finkelstein Y, Shapiro-Feinberg M, Stein G, Ophir D. Uvulopalatopharyngoplasty vs. laser-assisted uvulopalatoplasty. Arch Otolaryngol Head Neck Surg 1997;123:265-76.
- 26. Maw J. Uvulopalatopharyngoplasty versus laser-assisted uvulopalatopharyngoplasty in the treatment of snoring. J Otolaryngol 1997;26(4):232-5.
- 27. Wennmo C, Olsson P, Flisberg K, Paulsson B, Luttrup S. Treatment of snoring—with and without carbon dioxide laser. Acta Otolaryngol (Stockh) 1992;Suppl 492:152-5.
- 28. Hagert B, Wahren LK, Wikblad K, Ödkvist L. Patients' and cohabitants' reports on snoring and daytime sleepiness, 1-8 years after surgical treatment of snoring. ORL 1999;61:19-24.
- 29. Coleman JA. Laser-assisted uvulopalatoplasty: long-term results with a treatment for snoring. ENT Journal 1998;77(1):23-34.
- 30. Wareing MJ, Callanan VP, Mitchell DB. Laser assisted uvulopalatoplasty: six and eighteen month results. J Laryngol Otol 1998;112:639-41.
- 31. Finkelstein Y, Ophir D. One-stage laser-assisted uvulopalatoplasty [letter to the editor]. J Otolaryngol 1997;26(2):147.
- 32. Troell RJ, Powell NB, Riley RW, Li KK, Guilleminault C. Comparison of postoperative pain between laser-assisted uvulopalatoplasty, uvulopalatopharyngoplasty, and radiofrequency volumetric tissue reduction of the palate. Otolaryngol Head Neck Surg 2000;122:402-9.
- 33. Walker RP, Gopalsami C. Laser-assisted uvulopalatoplasty: postoperative complications. Laryngoscope 1996;106:834-8.
- 34. Pinczower EF. Globus sensation after laser-assisted uvuloplasty. Am J Otolaryngol 1998;19(2):107-8.
- 35. Esberg A, Levring-Jäghagen E, Dahlström M, Dahlqvist Å. Persistent dysphagia after laser uvulopalatoplasty. Acta Otolaryngol (Stockh) 1998;118:870-4.
- 36. Issa FG, Sullivan CE. Upper airway closing pressures in snorers. J Appl Physiol 1984;57(2):528-35.
- 37. Baggish MS, Elbakry M. The effects of laser smoke on the lungs of rats. Am J Obstet Gynecol 1987;156(5):1260-5.
- 38. Berger G, Finkelstein Y, Ophir D. Histopathologic changes of the soft palate after laser-assisted uvulopalatoplasty. Arch Otolaryngl Head Neck Surg 1999;125:786-90.
- 39. Sackett D. Rules of evidence and clinical recommendation. Can J Cardiol 1993;9:487-9.
- 40. Kales A, Cadieux RJ, Bixler EO, Soldatos CR, Vela-Bueno A, Misoul CA, Locke TW. Severe obstructive sleep apnea-I: Onset, clinical course, and characteristics. J Chronic Dis 1985;38(5):419-25.
- 41. Eddy DM, eds. A manual for assessing health practices and designing practice policies: The explicit approach. Philadelphia, PA: American College of Physicians, 1992.
- 42. Pinczower EF. Globus sensation after laser-assisted uvulopalatoplasty. Am J Otolaryngol 1998;19(2):107-8.

- 43. Cheng D, Weng J, Yang P-W. Carbon dioxide laser surgery for snoring: results in 192 patients. Otolaryngol Head Neck Surg 1998;118:486-9
- 44. O'Reilly BF, Simpson DC. A comparison of conservative, radical and laser palatal surgery for snoring. J R Coll Surg Edinb 1998;43:194-5.
- 45. Isberg A, Levring-Jäghagen E, Dahlström M, Dahlqvist Å. Persistent dysphagia after laser uvulopalatoplasty. Acta Otolaryngol (Stockh) 1998;118:870-4.
- 46. Ingrams DR, Spraggs PDR, Pringle MB, Croft CB. CO2 laser palatoplasty: early results. J Laryngol Otol 1996;110:754-6.
- 47. Wareing M, Mitchell D. Laser-assisted uvulopalatoplasty: an assessment of a technique. J Laryngol Otol 1996;110:232-6.
- 48. Pribtikin EA, Schutte SL, Keane WM, Mao V, Cater JR, Doghramji K, Youakim JM, Rosen MR, Breuninger W. Efficacy of laser-assisted uvulopalatoplasty in obstructive sleep apnea. Ototlaryngol Head Neck Surg 1998;119:643-7.
- 49. Kotecha B, Paun S, Leong P, Croft CB. Laser assisted uvulopalatoplasty: an objective evaluation of the technique and results. Clin Otolaryngol 1998;23:354-9.
- 50. Skatvedt O. Laser-assisted uvulopalatoplasty: description of the technique and pre- and postoperative evaluation of subjective symptoms. ORL 1996;58:243-7.
- 51. Gnuechtel MM, Keyser JS, Greinwald, JR., JH, Postma GN. Electrocautery versus carbon dioxide laser for uvulopalatoplasty in the treatment of snoring. Laryngoscope 1997;107:848-54.
- 52. Krespi YP, Khosh MM. The efficacy of laser-assisted uvulopalatoplasty in the management of obstructive sleep apnea. In Krespi YP, ed., Office-based surgery of the head and neck. Philadelphia, PA: Lippincott-Raven, 1998:129-140.
- 53. Kamami Y-V. Outpatient treatment of sleep apnea syndrome with CO₂ laser: laser-assisted UPPP. J Otolarynogol 1994;23:395-8.
- 54. Coleman J, Rathfoot MC. Oropharyngeal surgery in the management of upper airway obstruction during sleep. Otolaryngologic Clin N Amer 1999;32(2):263-76.
- 55. Ikeda K, Oshima T, Tanno N, Ogura M, Shimomura A, Suzuki H, Takasaka T. Laser-assisted uvulopalatoplasty for habitual snoring without sleep apnea: outcome and complications. ORL 1997;59:45-9.
- 56. Kamami Y-V. Outpatient treatment of snoring with CO2 laser: laser-assisted UPPP. J Otolaryngol 1994;23:391-4.
- 57. Carenfelt C, Haraldsson PO. Frequency of complications after uvulopalatopharygoplasty. [letter to the editor] Lancet 1993; 341(8842):437.

Table 1. LAUP Controlled Trials

LAUP, laser-assisted uvulopalatoplasty; DVT, deep venous thrombosis; NPO, nocturnal pulse oximetry; NRan, nonrandomized; Ran, randomized; *, significant difference; UPPP, uvulopalatopharyngoplasty; VPI, velopharyngeal insufficiency; PSG, polysomnography; AHI, apnea/hypopnea index; RDI, respiratory disturbance index which is interchangeable with AHI

Evidence	Reference/		
Procedure: Number of			
Sample Size			
Outcome			
	-		

Finkelstein (25) Level III -C	Walker (24) Level III -C		Remacle (3) Level III -C	Reference/ Evidence Level
NRan	NRan		NRan concurrent cohort study	Study Design
First 100/174 consecutive patients had UPPP, remaining 74/174 had LAUP (first 34/74 had incision comparable to UPPP with	LAUP vs. UPPP; 1 or more LAUP procedures; UPPP included tonsillectomy (32/41) and nasal surgery (25/41); postop PSG after 3 mo	postop PSG after 6 mo	LAUP vs. UPPP; decisional algorithm: LAUP or UPPP if AHI <40 or if AHI >40 and fails CPAP; UPPP if hypertrophic palatine tonsils and long or	Procedure; Number of Sessions; Protocol
Snoring: negative for OSA by PSG OSA defined as PSG with RDI >5	OSA: PSG with RDI >5		Snoring: patient history OSA: screening NPO; if positive, PSG	Diagnostic Criteria
N= 174 / 22- 71 y / 157M, 17F (16 with heavy snoring only)	N=167, 79 completed (38 LAUP / 31M, 7F / mean = 53.6 y; 41 UPPP / 40M, 1F / mean = 45.7 y)	for OSA)	N=89 / 23- 77 y / 70M, 19F; 78 completed (63 had surgery for habitual	Sample Size / Age / Sex of Subjects
Intraoral photographs of soft palate; peroral and nasopharyn- goscopic examination	PSG: >50% RDI reduction in postop PSG vs. preop PSG	PSG: normal, AHI <10; improved, AHI decreased by at least 1 AHI stage (>10 and <20, >20 and <40, >40); failure, no change in AHI stage	Questionnaires NPO: normal, <90% in <1% of night and <5% of desat of mean O ₂ sat; improved, 1 of 2 criteria met; fell to the content of the	Outcome Measures
None reported	LAUP: 2/38 bleeding; 2/38 oral candidiasis; 1/38 temp VPI UPPP: 2/41 bleeding; 3/41 temp VPI; 1/41 lower extremity DVT	dysphonia x 1 y	LAUP: 6% minor dysphonia x 1y; 1 case severe dysphagia; 1 case minor bleeding UPPP: 20% temp nasal regurgitation;	Adverse Effects
LAUP: circumferential scarring, resulting in decreased velopharyngeal air space and decreased	Postop RDI >50% reduction: 18/38 LAUP; 21/41 UPPP Postop RDI reduction: 30.3 to 22.2 LAUP*; 52.1 to 25.5 UPPP* Postop min O ₂ change: 83.3 to 81.6%; LAUP; 72.8 to 80.9% UPPP*	8.60, resp) 8.60, resp) NPO: postop 8 normal, 5 improved, 1 failure, 1 refused PSG: postop 4 normal, 2 improved, 2 failures, 7 refused	LAUP vs. UPPP: no sig difference in pain level or duration by questionnaires, no sig difference in satisfaction on 0-10	Conclusions
			Significant dropout rate for return PSG (7/15 refused follow-up PSG)	Comments

Wennmo (27) Level III -C	Maw (26) Level III -C		Reference/ Evidence Level
NRan	NRan		Study Design
LAUP vs. UPPP vs. UPPP with tonsillectomy; patients in the first 2 groups were selected for small tonsils; follow-up from 3 mo to 2 y	LAUP vs. tonsillectomy and UPPP; eligible if Müller maneuver showed 75% and ≤50% obstruction at level of soft palate and tongue base, resp; ≥50% obstruction due to tonsillar hypertrophy = UPPP, <50% = LAUP (up to 4 stages); follow-up 4 wk postop	tonsillotomy; remaining 40/74 uvula reduced); follow-up intraoral photography, peroral and nasopharyngoscopic examination up to 12 weeks postop	Procedure; Number of Sessions; Protocol
Snoring: questionnaire	Snoring: history, questionnaire, PSG with AHI ≤20		Diagnostic Criteria
N=30 (10 LAUP / mean = 47 y / M:F ratio = 9:1; 10 UPPP / mean = 45.3 y / M:F ratio = 10:0; 10 UPPP with tonsillectom y / mean = 44.5 y / M:F ratio = 9:1)	N=136; 129 completed (80 LAUP / mean = 50 y / 88% M; 29 UPPP / mean = 41 y / 93% M)		Sample Size / Age / Sex of Subjects
Questionnaire	Questionnaire		Outcome Measures
UPPP with tonsillectomy: 1 with bleeding; 2 with minor oropharyngeal discomfort	LAUP: pain level equivalent to UPPP; 3/80 delayed postop bleeding UPPP: 1/29 delayed postop bleeding		Adverse Effects
Snoring: subjective improvement in all patients except for one in the UPPP with tonsillectomy group	LAUP vs. UPPP: no sig differences for final snoring scores, pain, or complication rates. Selection by Müller maneuver allowed >50% reduction in final snoring scores in 97% of patients.	distensibility UPPP: enlarged oropharynx and increased velopharyngeal air space	Conclusions
Small sample size; selection bias			Comments

Carenteit (21) Level III -C	Reference/ Evidence Level
N Aan	Study Design
randomized 33 patients for LAUP and 37 patients for UPPP, then added an additional 30 consecutive patients for LAUP; follow-up 3-4 mo postop	Procedure; Number of Sessions; Protocol
at level of velopharynx, distance between faucial tonsils >25 mm, no simultaneous nasal surgeries performed, PSG <10 obstructive apneas >10 sec	Diagnostic Criteria
N=100 (63 LAUP/24- 70 y / 18% F; 37 UPPP / 30-74 y / 14% F); returned for follow-up: 60 LAUP; 36 UPPP	Sample Size / Age / Sex of Subjects
Questionnaire	Outcome Measures
with incomplete surgery due to strong vomiting reflexes; 16 patients with slight or moderate scar fibrosis; 2 patients with scar fibrosis with narrowed nasophayngeal aperture and nasal obstruction, worsened scarring after reoperation with UPPP UPPP: 5 patients with slight or moderate scar fibrosis	Adverse Effects
lotal or near-total snoring elimination: 51/60 LAUP; 32/36 UPPP Without complaints from family: 56/60 LAUP; 33/36 UPPP No habitual daytime sleep attacks: 15/21 LAUP; 10/13 UPPP	Conclusions
Considered a nonrandomized study due to selection bias and partial randomization of LAUP group	Comments

Table 2. LAUP OSA Case Series and Cohort Studies

LAUP, laser-assisted uvulopalatoplasty; LSAT, lowest oxygen saturation; *, significant difference; UPPP, uvulopalatopharyngoplasty; MPS, multilevel pharyngeal surgery; STR, septoplasty with turbinate reduction; VPI, velopharyngeal insufficiency; PSG, polysomnography; AHI, apnea/hypopnea index; RDI, respiratory disturbance index which is interchangeable with AHI; UARS, upper airway resistance syndrome; MSLT, multiple sleep latency test; HA, headache

	Walker (13) Level V		Ryan (12) Level V	Reference/ Evidence Level
	Case Series		Case Series	Study Design
	LAUP; postop PSG after 3 mo, long-term questionmaire data		LAUP; one-stage resection; postop PSG after 3 mo	Procedure; Number of Sessions; Protocol
	OSA by PSG		OSA by PSG	Diagnostic Criteria
data and without interim surgery or PSG <6 wks postop, 31 with long-term data / 35-75 y / 30M, 10F	N=182, 131 completed treatment, 40 with complete poston PSG		N=44 / mean = 49 y / 37M, 7F	Sample Size / Age / Sex of Subjects
	PSG Questionnaire	poor response, AHI >50% of pre-LAUP value; worse, AHI >100% of pre- LAUP value Videoendo- scopy Questionnaires	PSG: good response, AHI ≤10; partial response, AHI ≤50% of pre- LAUP value;	Outcome Measures
	None reported	\$	excessive mouth dryness, throat pain or discomfort	Adverse Effects
17.6 Questionniare: presenting complaints improved in 74.9% after mean followup of 4.04 y in 31 subjects	PSG: AHI improved* from mean of 25.0 to 15.3, REM% increased* from mean of 13.3 to	sectional and anteroposterior diameter* Questionnaires: improved quality of life*, sleepiness*, and snoring index	PSG: postop 12 good response, 4 partial response, 15 poor response, 13 worse Videoendo-scopy:	Conclusions
				Comments

Mickelson (14) Level V	Reference/ Evidence Level
Case Series	Study Design
LAUP; PSG 6-12 wks postop, questionnaire data from patient and bedpartner before LAUP, 6-12 wks postop, and >2 y	Procedure; Number of Sessions; Protocol
PSG: OSA defined as RDI >10	Diagnostic Criteria
N= 59, 36 completed postop PSG; MSLT pre- and postop in 7 patients / mean = 52.3 y/ 29M, 7F	Sample Size / Age / Sex of Subjects
PSG MSLT Questionnaires	Outcome Measures
I patient with several drops of bleeding controlled with cautery; 1 patient taking ibuprofen for pain who bled about 30 ml on postop day 4 that ceased spontaneously	Adverse Effects
PSG: AHI decreased* from mean of 28.1 to 7.9, min O2 saturation increased from mean of 80.6% to 84.0% MSLT: mean sleep latency improved* Questionnaire: improved* snoring, morning fatigue, morning HA, daytime somnolence, daytime psychometric measures	Conclusions
	Comments

Walker (15) Level V	Reference/ Evidence Level
Case Series	Study Design
LAUP; 3-6 treatments; postop PSG≥ 3 mos after LAUP	Procedure; Number of Sessions; Protocol
PSG: mild OSA (AHI >5 and <20), moderate OSA (AHI >20 and <39), severe OSA (AHI ≥40)	Diagnostic Criteria
N= 38/39- 75 y/31M, 7F	Sample Size / Age / Sex of Subjects
PSG: surgical response rate defined as >50% AHI reduction in postop PSG vs. preop PSG and a postop AHI<20	Outcome Measures
2 with bleeding, 2 with oral candidiasis, I with temporary VPI	Adverse Effects
PSG: mild OSA AHI decreased from 10.5 to 10.4 (surgical response rate of 46.7%) and LSAT decreased from 87.2% to 86.8%; moderate OSA AHI decreased from 29.0 to 21.1 (surgical response rate of 41.7%) and LSAT decreased from 81.3% to 80.4%; severe OSA AHI decreased from 59.7 to 39.6 (surgical response rate of 45.5%) and LSAT increased from 80.3% to 81.1%	Conclusions
	Comments

Utley (16) Level V	Reference/ Evidence Level
Retro- spective Cohort	Study Design
LAUP; mild OSA (AHI >5 and <20) patients encouraged to undergo LAUP or UPPP, if they had significant findings on the modified Muller maneuver, they were encouraged to have MPS; those patients with moderate OSA (AHI ≥20 and <40) or severe OSA (AHI ≥40) were offered MPS; those with significant nasal obstruction unresponsive to medication were offered STR; postop PSG min of 4 mos after LAUP	Procedure; Number of Sessions; Protocol
Snoring UARS OSA: surgical response rate defined as >50% drop in AI or AHI, with a postop AI <10 or postop AHI <20	Diagnostic Criteria
N= 229, 95 candidates for surgery, 56 for LAUP (12 with postop PSG) and 32 for MPS (14 with postop PSG) and 6 for STR/ LAUP mean = 45.3 y, MPS mean = 48.8 y/ LAUP 50M, 6F, MPS 30M 2F, STR 5M 1F	Sample Size / Age / Sex of Subjects
PSG Modified Muller maneuver Questionnaires	Outcome Measures
LAUP: 1 with vasovagal episode during LAUP, 1 with bleeding requiring electrocautery MPS: all with transient paresthesia of mandibular incisors, 5 with gingivolabial sulcus incision dehiscences, 2 extruded screws and mandibular bony segments, 1 with moderate ecchymosis and edema of the neck and face skin without airway compromise, 1 with gingivolabial sulcus wound infection STR: none reported	Adverse Effects
PSG: LAUP OSA surgical response rate of 41.7% (5/12), MPS response rate of 85.7% (12/14), STR 16.7% (1/6) Modified Muller maneuver: palatal collapse decreased* post LAUP; collapse at all 3 levels decreased* post MPS decreased* post LAUP*, post MPS Epworth: improved post LAUP*, post MPS Epworth: complete cure in 41.8% post LAUP, 50% po	Conclusions
pre- and postop PSG were different mixtures of attended and unattended studies; some patients in LAUP group had UARS	Comments

		Level V	Evidence Level Skatvedt (5)
			Study Design Case Series
			Procedure; Number of Sessions; Protocol LAUP; postop PSG 3- 16 mos after LAUP
			Diagnostic Criteria OSA by PSG
		-	Sample Size / Age / Sex of Subjects N= 16 / 26- 63 y / 15M,
		pharyngeal and esophageal pressure measures with a nasal tube containing 6 pressure sensors Questionnaire	Outcome Measures PSG Continuous
			Adverse Effects None reported
nypopneas, 92% preop(85% postop Questionnaire: reduced* incidence and loudness of snoring, apneas, morning fatigue, excessive daytime sleepiness, and morning HA	ANKEM time associated with O2 sats <80% Continuous pressure measures: velopharyngeal obstructive segments in 90% preop (9% postop), and in	respiratory events, AHI, incidence of sleep with snoring, microarousal index, and mean duration of NREM sleep related to total	Conclusions PSG: improved* duration of
			Comments

					-				2	*		Level V	(17)	Walker		Level	Evidence	Reference/
								-						Case Series	Design	Study		
										after LAUP	postop PSG ≥ 3 mos	treatments for OSA;	for snoring, 1-7	LAUP; 1-5 treatments		Sessions; Protocol	Procedure; Number of	
									-			አ ·	defined as RDI	PSG: OSA	Criteria	Diagnostic		
		•				-	28M, 5F	=51.9 y/	PSG) / mean	with postop	OSA (33	65 with	with snoring,	N= 170, 105	of Subjects	/ Age / Sex	Sample Size	
The second secon			-							Questionnaires	RDI ≤10	defined as postop	surgical success	PSG: OSA		Measures	Outcome	
			-											None reported		Adverse Effects		
	change	without significant	results, 15%	21% with worse	surgical success;	OSA: 48% with	improvement	without	improvement, 10%	partial	snoring, 29% with	elimination of	complete	Snoring: 60% with		Conclusions		
																Comments		

Table 3. Effect Size of LAUP OSAS Case Series and Cohort Studies

Reference	Number of	AHI Mean (SD)		D*	T-654 C:**	0
Reference	subjects	Pre-LAUP	Post-LAUP	Duration*	Effect Size**	Comments
Ryan (12)	44	29 (17)	19 (15)	min 3 mos	0.588	Calculated SD from SE
Walker (13)	40	25 (17.7)	15.3 (18.3)	48-896 days	0.548	Calculated SD from SE
Mickelson (14)	36	28.1 (17.3)	17.9 (13.5)	6-12 weeks	0.590	
Walker (15)	38	30.6 (22.6)	22.2 (26.8)	min 3 mos	0.369	
Utley (16)	12	8.9 (6.1)	10.3 (8.1)	min 4 mos	-0.230	
Skatvedt (5)	16	18.6 (23.4)	6.4 (10.2)	3-16 mos	0.522	
Walker (17)	33	29.4 (21.2)	21.8 (24.7)	min 3 mos	0.358	
Summary	219	NA	NA	NA	0.392	Unadjusted average of effect sizes***
Summary	219	NA	NA	NA	0.251	Adjusted average of effect sizes****

SD - Standard Deviation of the mean

SE - Standard error of the mean

SE = SD/(square root of the number of subjects)

NA - Not applicable

Table 4. AASM Classification of Evidence

Recommendation	Evidence	Study
Grades	Levels	Design
A	I	Randomized well-designed trials with low-alpha & low-beta errors*
В	II	Randomized trials with high-beta errors*
С	III	Nonrandomized controlled or concurrent cohort studies
С	IV	Nonrandomized historical cohort studies
C	V	Case series

ADAPTED FROM SACKETT 39

Table 5. AASM Levels of Recommendations

Term	Definition
Standard	This is a generally accepted patient-care strategy, which reflects a high degree of clinical
	certainty. The term standard generally implies the use of Level I Evidence, which directly addresses the clinical issue, or overwhelming Level II Evidence.
Guideline	This is a patient-care strategy, which reflects a moderate degree of clinical certainty. The
	term guideline implies the use of Level II Evidence or a consensus of Level III Evidence
Option	This is a patient-care strategy, which reflects uncertain clinical use. The term option implies
	either inconclusive or conflicting evidence or conflicting expert opinion.

ADAPTED FROM EDDY 41

^{*} Time from last LAUP treatment to post-LAUP PSG

^{**} Effect size = (Pre-LAUP AHI mean - Post-LAUP AHI mean) / Pre-LAUP AHI standard deviation

^{***}unadjusted average is the sum of the individual effects sizes/the number of studies (7 in this case)

^{****}adjusted average is 1/SE² times [(Pre-LAUP AHI mean - Post-LAUP AHI mean)/ Pre-LAUP AHI standard deviation] for each study and summing the results. This sum is divided by the sum of the 1/SE² for each of the 7 studies where SE is the standard error of the pre-LAUP mean

^{*}Alpha error refers to the probability (generally set at 95% or greater) that a significant result (e.g., p<0.05) is the correct conclusion of the study or studies. Beta error refers to the probability (generally set at 80% or 90% or greater) that a nonsignificant result (e.g., p>0.05) is the correct conclusion of the study or studies. The estimation of beta error is generally the result of a power analysis. The power analysis includes a sample size analysis which projects the size of the study population necessary to ensure that significant differences will be observed if actually present.

Table 6. LAUP Adverse Effects

Adverse Effects*	Frequency (%)	References
Choking at meals	81	42
Dysphagia		
Temporary	31	43
Persistent	5 - 53	44,45
Severe	1	3
Poor appetite	21	43
Dry throat (Persistent)	16 - 42	44,46,47
Problems drinking (Persistent)	16	44
Globus sensation (Persistent)	10 - 25	42,45,47,48
Increased gag reflex	10	48
Differences in swallowing (Persistent)	6	47
Dysphonia (Mild)	6	3
Vasovagal episode	1.8	16
Voice change (Temporary)	1.7 - 17.2	46,49
Nasal regurgitation		
Temporary	1.7 - 10.3	43,46,49
Persistent	1 - 20	44,45,47,50
Vomiting	1.5	21
Bleeding		
Non-severe, immediate or delayed postoperative, includes	1 - 8	3,15,24,26,33,43,49,51,52,53,54,55
hemoptysis		
Requiring medical attention	0.4 – 1.8	16,23,33,54
Velopharyngeal insufficiency (Temporary)	0.5 - 3	24,33,44,51,56
Loss of taste		
Temporary	0.3	33
Persistent	5	48
Scar fibrosis	0.2 – 30 (100**)	21,38,56
Mild-Moderate	25	21
Severe	3	21
Infection		
Bacterial	0.13	33
Oral candidiasis	0.4 - 5.3	24,33,52
Septicemia - fatal	.03 (1/2900)	57
Indeterminate type	0.4 - 2	23,51

^{*} Other than temporary postoperative pain not exceeding 3 weeks in duration; Temporary defined as equal to or less than one month duration; Persistent = greater than one month duration; Reported nonspecific or vague symptomatology are not included.

** Histopathologic study