Washington University School of Medicine Digital Commons@Becker

2020-Current year OA Pubs

Open Access Publications

6-1-2022

A guide for managing patients with stage I NSCLC: Deciding between lobectomy, segmentectomy, wedge, SBRT and ablationpart 2: Systematic review of evidence regarding resection extent in generally healthy patients

Frank C. Detterbeck Vincent J. Mase Andrew X. Li Ulas Kumbasar

Brett C. Bade

See next page for additional authors

Follow this and additional works at: https://digitalcommons.wustl.edu/oa_4

Authors

Frank C. Detterbeck, Vincent J. Mase, Andrew X. Li, Ulas Kumbasar, Brett C. Bade, Henry S. Park, Roy H. Decker, David C. Madoff, Gavitt A. Woodard, Whitney S Brandt, and Justin D. Blasberg



A guide for managing patients with stage I NSCLC: deciding between lobectomy, segmentectomy, wedge, SBRT and ablation – part 2: systematic review of evidence regarding resection extent in generally healthy patients

Frank C. Detterbeck¹[^], Vincent J. Mase Jr¹, Andrew X. Li², Ulas Kumbasar³, Brett C. Bade⁴, Henry S. Park⁵, Roy H. Decker⁵, David C. Madoff⁶, Gavitt A. Woodard¹, Whitney S. Brandt⁷, Justin D. Blasberg¹

¹Department of Thoracic Surgery, Yale University School of Medicine, New Haven, CT, USA; ²Department of General Surgery, Yale University School of Medicine, New Haven, CT, USA; ³Department of Thoracic Surgery, Hacettepe University School of Medicine, Ankara, Turkey; ⁴Department of Pulmonary Medicine, Yale University School of Medicine, New Haven, CT, USA; ⁵Department of Therapeutic Radiology, Yale University School of Medicine, New Haven, CT, USA; ⁶Department of Radiology & Biomedical Imaging, Yale University School of Medicine, New Haven, CT, USA; ⁷Department of Cardiothoracic Surgery, Washington University School of Medicine, St. Louis, MO, USA

Contributions: (I) Conception and design: FC Detterbeck; (II) Administrative support: None; (III) Provision of study materials or patients: None; (IV) Collection and assembly of data: All authors; (V) Data analysis and interpretation: All authors; (VI) Manuscript writing: All authors; (VII) Final approval of manuscript: All authors.

Correspondence to: Frank C. Detterbeck, MD. Professor of Thoracic Surgery, Yale University School of Medicine, P.O. Box 208062, New Haven, CT, USA. Email: frank.detterbeck@yale.edu.

Background: Clinical decision-making for patients with stage I lung cancer is complex. It involves multiple options (lobectomy, segmentectomy, wedge, stereotactic body radiotherapy, thermal ablation), weighing multiple outcomes (e.g., short-, intermediate-, long-term) and multiple aspects of each (e.g., magnitude of a difference, the degree of confidence in the evidence, and the applicability to the patient and setting at hand). A structure is needed to summarize the relevant evidence for an individual patient and to identify which outcomes have the greatest impact on the decision-making.

Methods: A PubMed systematic review from 2000–2021 of outcomes after lobectomy, segmentectomy and wedge resection in generally healthy patients is the focus of this paper. Evidence was abstracted from randomized trials and non-randomized comparisons with at least some adjustment for confounders. The analysis involved careful assessment, including characteristics of patients, settings, residual confounding etc. to expose degrees of uncertainty and applicability to individual patients. Evidence is summarized that provides an at-a-glance overall impression as well as the ability to delve into layers of details of the patients, settings and treatments involved.

Results: In healthy patients there is no short-term benefit to sublobar resection *vs.* lobectomy in randomized and non-randomized comparisons. A detriment in long-term outcomes is demonstrated by adjusted non-randomized comparisons, more marked for wedge than segmentectomy. Quality-of-life data is confounded by the use of video-assisted approaches; evidence suggests the approach has more impact than the resection extent. Differences in pulmonary function tests by resection extent are not clinically meaningful in healthy patients, especially for multi-segmentectomy *vs.* lobectomy. The margin distance is associated with the risk of recurrence.

Conclusions: A systematic, comprehensive summary of evidence regarding resection extent in healthy patients with attention to aspects of applicability, uncertainty and effect modifiers provides a foundation on

^ ORCID: 0000-0002-2858-8016.

2358

which to build a framework for individualized clinical decision-making.

Keywords: Lung cancer; surgery; lobectomy; segmentectomy; wedge

Submitted Nov 19, 2021. Accepted for publication May 05, 2022. doi: 10.21037/jtd-21-1824 View this article at: https://dx.doi.org/10.21037/jtd-21-1824

Introduction

Treatment options for clinical stage I (cI) non-small cell lung cancer (NSCLC) have evolved. Smaller tumors are being detected; average patient age is increasing, as is the number with co-morbidities. We need to match the treatment to the patient and tumor, avoiding both overtreatment and undertreatment.

Decision-making regarding stage I NSCLC is complex. Many short- and long-term outcomes are relevant. We aim to practice evidence-based medicine (EBM), but the available evidence is suboptimal and confusing. Multiple factors influence treatment selection and independently the prognosis, and evidence often only partially applies to an individual patient. Although clinicians are used to weighing various considerations and complex decision-making, better definition of the evidence regarding management of cI NSCLC is needed, including sources of uncertainty, and nuances of patients, tumors and settings that affect applicability.

We assessed the evidence regarding cI NSCLC, critically addressing confounders and limitations, to provide clarity and confidence in applicability in various circumstances. Furthermore, we developed a concise format that enhances application to individual patients. The project consists of 4 publications: Part 1 concisely summarizes the evidence and provides a framework to guide clinical decision-making (1), Part 2 (this paper) reviews evidence regarding surgery in generally healthy patients, Part 3 addresses surgery in specific patients and tumors (2), Part 4 focuses on evidence regarding SBRT and ablation (3).

Methods

General approach

The approach involved being as inclusive and as critical as possible, with attention to nuances about settings and characteristics of the available evidence to understand limitations and applicability. A detailed description of the approach is provided in the methods section of Part 1 (1). Briefly, the subject is stage cIA NSCLC (using the 8th edition nomenclature throughout); interventions include lobectomy, segmentectomy, wedge resection, SBRT and ablation. The most relevant outcomes were chosen *a priori*: short-term treatment-related mortality, toxicity/morbidity, pain, quality-of-life (QOL) and longterm overall survival (OS), lung cancer specific survival (LCSS), freedom from recurrence (FFR), functional status and QOL.

Because few randomized controlled trials (RCTs) are available, we relied heavily on non-randomized comparisons (NRCs) that adjusted for confounding factors (i.e., factors independently influencing treatment selection and outcomes). We critically evaluated how well confounders were accounted for to assess the confidence that observed results reflect the intervention in question. Finally, we explored sources of ambiguity to promote understanding uncertainties and limitations of applicability.

Clinical decision-making requires weighing multiple considerations for an individual. This involves balancing not only many outcomes but many aspects of each—e.g., the strength of the evidence, the magnitude of the impact, uncertainty and how well this applies to an individual. In the Part 1 paper we provide a framework to manage this complexity—allowing clinicians to identify and focus on issues with the most impact in a particular setting for a patient. Here we develop the foundation, presenting the data in a manner that can at-a-glance provide an aggregate view of an outcome as well as the nuances and uncertainties of the data. A definition of what can be reasonably considered clinically meaningful facilitates assessing the impact of differences (described elsewhere; see *Tab. S1-1* of Part 1) (1).

Evidence assessment

Literature search and study selection

We systematically searched English literature from 2000-2021; details are provided elsewhere (see *app. 1-2* of Part 1) (1). Selected studies provided evidence

Journal of Thoracic Disease, Vol 14, No 6 June 2022

relevant to the topic, focusing on RCTs and adjusted NRCs. For major outcomes we included all RCTs, and NRCs that adjusted for confounding and had \geq 50 patients per arm. Each evidence table lists specific inclusion and exclusion criteria.

Study assessment

NRCs were assessed for confounding (bias) in order to appropriately interpret findings. The assessment of NRCs is summarized below (details provided in Appendix 2-1).

Potential confounders

A comprehensive list of potential confounders was identified *a priori* from known prognostic factors, patterns of care and treatment discrepancies. These included non-medical patient-related factors (e.g., age, sex, race, education, socioeconomic, marital status), medical patient-related factors [e.g., comorbidities, comorbidity severity, performance status (PS)], discrepancies in stage classification [e.g., node assessment, positron emission tomography (PET) use], time period (treatments skewed towards different periods), facility factors (treatments skewed towards different facility types), treatment quality (e.g., margin adequacy, experience, technical aspects), favorable tumor selection [e.g., smaller, ground glass (GG), indolent tumors, conversion to lobectomy if upstaging suspected/encountered].

Methods of multivariable adjustment

Multivariable regression models the relationship between multiple covariates and an outcome. Simultaneous adjustment for multiple confounders requires a substantial sample size—generally ~10 events (e.g., deaths) for each covariate. Propensity scoring models the relationship between confounders and treatment assignment, collapsing all confounders into a single propensity score. While theoretically advantageous when there are many confounders and few events, whether propensity or multivariable methods more accurately estimate treatment effect is unclear (4,5). Several propensity adjustment methods exist (propensity score adjustment, matching, inverse weighting); performance of each depends on characteristics of the data and question at hand (4-6).

Assessment of confidence study results reflect the treatment of interest

Relevant NRCs were assessed using a general tool to assess overall risk of bias (7). Additionally, we developed an assessment specific to stage I lung cancer, based on the *a priori* list of potential confounders (details in Appendix 2-1). Two reviewers rated each domain in each study and intervention, assigning an overall degree of confidence that outcomes reflect the treatment intervention; discrepancies were resolved by discussion. The independent assessments were largely consistent (and similar to the general tool rating), providing confidence in the process. The evidence tables include the consensus ratings for residual confounding.

Aggregation of studies

A quantitative meta-analysis is deemed inappropriate because of frequent residual confounding in various domains with variable severity. It is more useful to aggregate the studies in a manner that highlights similarities and differences, with ordering that allows patterns to emerge. This facilitates an overall qualitative impression that is more conducive to guiding clinical decision-making.

To achieve this, we have thoughtfully constructed tables. Color coding rapidly provides an overall impression (despite inclusion of levels of details if close scrutiny is needed). This essentially layers the concept of a heat map onto a traditional table. We explored various ways of ordering table entries, eventually settling on what was most revealing regarding the presence/absence of an association. The table structure is noted as a subtitle. We believe that visual representation of the outcomes, uncertainties and effect modifiers provides a summary that enhances point-of-care clinical judgment.

Results

Sbort-term outcomes

Treatment related mortality

Several RCTs reveal no difference in mortality by resection extent in healthy patients. The Lung Cancer Study Group (LCSG821) trial, conducted in the 1980s, reported no significant mortality difference between sublobar resection (2/3rd segmentectomy) and lobectomy via thoracotomy (1% vs. 2% respectively) (8). In a US-based RCT (CALGB140503, 2007–17) 90-day mortality was not statistically different for sublobar resection vs. lobectomy (1.2% vs. 1.7%; 80% VATS resection, 60% wedge among sublobar resection) (9). No mortality occurred for either segmentectomy or lobectomy in a large Japanese RCT (JCOG0802, 2009–14, n=1,106) (10) and a smaller European RCT (n=108) (11).

Studies of perioperative mortality with adjustment for

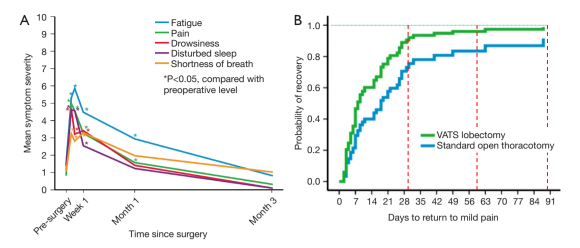


Figure 1 (A,B) Symptoms and recovery after lung resection.

Prospective study of patient reported outcomes in patients undergoing lobectomy at MD Anderson (stage I, II NSCLC, 2004–08, n=60, 48% VATS). (A) Time course of the 5 most severe symptoms; 11-point scale from 0 (not present) to 10 (as bad as you can imagine). (B) Time to return to mild pain at 2 contiguous measurements. Reproduced with permission from Fagundes *et al.* (22). VATS, video-assisted thoracoscopic surgery.

confounders (Table S2-1) (12-18) have frequently reported minimally lower mortality after lesser resection, but the magnitude of the difference is not clinically meaningful. A difference of >1% was only noted in one study (wedge resection *vs.* lobectomy) in subgroups of thoracotomy and patients with a forced expiratory volume in 1 second (FEV1) of <60% (12).

Similar (unadjusted) mortality for lesser resection and lobectomy is reported in large database studies (e.g., 30-day mortality of 1.51%, 1.55% and 1.6%, P=0.87 for wedge resection, segmentectomy and lobectomy in an NCDB study [2003–11] (16); 90-day mortality 3.7% and 4% for sublobar resection and lobectomy in a SEER-Medicare study [2003–9] (19); 90-day mortality of 0.5%, 0.7% and 1.2% for wedge, segment and lobectomy, respectively, in a 2010 Japanese national study) (20). However, an Australian study reported unadjusted 90-day mortality of 4.5% and 2.6% for sublobar resection and lobectomy, respectively [2008–14] (21).

Treatment-related morbidity

Treatment-related morbidity is similar in large RCTs between sublobar resection and lobectomy in healthy patients (any morbidity, 51% vs. 54% CALGB, 51% vs. 48% JCOG0802; grade \geq 3 14% vs. 15% CALGB, 4.5% vs. 4.9% JCOG0802, each study using different grading definitions; and grade \geq 3 pulmonary complications, 7%

vs. 10% CALGB, 2.4% vs. 1.8% JCOG0802, respectively) (9,10). A nonsignificant trend towards lower grade \geq 3 complications in wedge vs. segmentectomy was seen in the CALGB study (11% vs. 19%, P=0.13) (9). The small European RCT also found no significant difference in overall 90-day morbidity (17% segmentectomy vs. 26% lobectomy, P= NS) (11).

Adjusted NRCs suggest slightly lower grade ≥ 3 complications after sublobar resection (Table S2-1, borderline clinically significant). The 90-day unadjusted grade ≥ 3 complication rate was low in the 2010 Japanese national experience (4.4% wedge, 7.1% segmentectomy, 8.7% lobectomy) (20).

Short-term pain, QOL

Few QOL studies have parsed results to sublobar resection, so extrapolation from general studies is required. Presumably most symptoms are incision-related—thus largely driven by the approach (VATS *vs.* open); resection extent can be mainly expected to impact dyspnea.

A prospective study shows that symptoms after lung resection mostly resolve within several months (*Figure 1A,1B*) (22). Similarly, QOL studies report the initial impairment in many domains is improved by 3–6 months (see subsequent QOL section)—especially after VATS resection. The impact of sublobar resection is unclear (studies are confounded by varying VATS use).

Journal of Thoracic Disease, Vol 14, No 6 June 2022

A small RCT reported on QOL over 12 months (2013– 17, n=108, closed after accruing 19% of the target) (11). Global QOL was significantly decreased at discharge and 6 weeks, returning to baseline by 3 months, with no difference between arms (segmentectomy vs. lobectomy). Interpretation is hampered because VATS was used for 23% of segmentectomies and 43% of lobectomies (P<0.03); furthermore, 44% of segmentectomies were arguably "lobelike" (i.e., left upper trisegmentectomy, lingulectomy, or basilar quadri-segmentectomy). Pain outcomes were similar for segmentectomy vs. lobectomy throughout, but worse than baseline in both arms even at 12 months. Dyspnea was worse than baseline throughout the follow-up year (somewhat less after segmentectomy than lobectomy) (11).

Many studies of lobectomy (including RCTs, adjusted NRCs) report better outcomes with VATS *vs.* thoracotomy [including lower operative mortality, fewer complications, shorter hospital length of stay (LOS) and less pain] (23). A recent RCT of lobectomy by VATS *vs.* anterolateral thoracotomy found less pain and less QOL reduction in the VATS arm; the QOL impact resolved in most patients by 6 (VATS) to 12 weeks (thoracotomy) (24).

VATS is also beneficial in sublobar resections. An extensively adjusted NRC found fewer complications with VATS (rated as "very high" confidence that outcomes reflect VATS vs. open approach to segmentectomy) (25). A retrospective comparison of VATS vs. open segmentectomy found fewer pulmonary complications and shorter LOS after VATS (n=193, 2000-13, mostly healthy, lobectomy eligible patients) (26). Another retrospective comparison of VATS vs. open segmentectomy (n=104 vs. 121) found that VATS was associated with fewer pulmonary complications (15% vs. 30%, P=0.012), shorter LOS (5 vs. 7 days, P<0.001), and statistically non-significant differences in overall complications (26% vs. 34%), major complications (6% vs. 12%) and operative mortality (0 vs. 1.7%), respectively (27).

Nomori *et al.* assessed pain, comparing segmentectomy via thoracotomy, segmentectomy via hybrid-VATS (VATS camera with mini-thoracotomy) and lobectomy via complete VATS (n=220, 2012-15) (28). Short-term pain was less after VATS/hybrid-VATS than thoracotomy, but similar for hybrid-VATS segmentectomy or VATS lobectomy. By 3 months pain had resolved equally in all groups, with <5% requiring any analgesics (28).

Nuances and sources of ambiguity

The type of segmentectomy may play a role: multivariable

analysis of a prospective study observed more grade ≥ 2 pulmonary complications following complex *vs*. simple segmentectomy (7.7% *vs*. 6.1%) (10). Complex segmentectomy was defined as requiring division of >1 intersegmental plane. However, another study found no difference in morbidity or mortality following complex (n=117) or simple (n=92) VATS segmentectomy (29).

Long-term outcomes

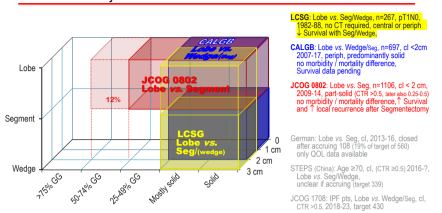
Survival

The LCSG821 RCT enrolled cN0 lung cancers ≤ 3 cm on the basis of CXR and not visible on (primarily rigid) bronchoscopy from 1982–88 (8,30,31). After intraoperative confirmation of T1N0 (frozen section of segmental, lobar, hilar, and mediastinal nodes)—patients were randomized to sublobar resection (67% segmentectomy) vs. lobectomy. A ≥ 2 cm margin was required; these tumors were undoubtedly primarily solid and resected via thoracotomy. In the final corrected analysis sublobar resection was associated with lower 5-year OS (56% vs. 73%; P=0.06), worse FFR (63% vs. 78%; P=0.04), and higher locoregional recurrence (5.4% vs. 1.9% per person per year, P=0.009) (30,31). However, present-day applicability of this evidence is questionable.

There are 2 major contemporary RCTs (*Figure 2*) (8-11,32-35). The CALGB140503 trial (9) randomized 697 patients with peripheral (outer 1/3), mostly solid tumors, ≤ 2 cm (total size) to sublobar resection (60% wedge) vs. lobectomy—mature results are awaited. The JCOG0802 trial (10,34) randomized 1,106 patients with peripheral (outer 1/3), part-solid tumors [88% with >0.5 consolidation/tumor ratio (CTR)], ≤ 2 cm (total size) to segmentectomy vs. lobectomy. A margin of ≥ 2 cm or a margin/tumor ratio ≥ 1 was required in both trials.

Long-term results of the JCOG0802 trial have been published (35), with similar results after segmentectomy *vs.* lobectomy. These results are discussed elsewhere (2) because this study involves part-solid tumors.

Adjusted NRCs of segmentectomy or wedge vs. lobectomy in apparently healthy patients are shown in *Table 1* (16,36-52), *Table 2* (16,36,42,47,48,50,53-62), *Table 3* (36,47,48,50,63-66) and Figures S2-1,S2-2,S2-3. Interpretation is challenging because of frequent limited accounting for confounders. Nevertheless, in aggregate, several observations can be made. First, the number of studies is impressive, and how inadequately most studies accounted for confounding factors. Second, the hazard ratios (HRs) for OS favor lobectomy (with few exceptions);



Major Randomized Controlled Trials

Figure 2 Major randomized controlled trials of lesser resection vs. lobectomy.

Graphic depiction of the 3 major randomized controlled trials. The x axis depicts the type of tumors included relative to proportion of solid/ground glass component, the z axis depicts tumor size, the y axis the resection extent. Three additional RCTs (German, STEPS and JCOG1706) are listed which have limited accrual. References: LCSG (8), CALGB (9), JCOG0802 (10), German (11), STEPS (32), JCOG1706 (33). CALGB, Cancer and Leukemia Group B; CTR, consolidation/tumor ratio; GG, ground glass appearance; IPF pts, Idiopathic pulmonary fibrosis patients; JCOG, Japan Cancer Oncology Group; LCSG, Lung Cancer Study Group; Lobe, lobectomy; Periph, peripheral; QOL, quality of life; Seg, segmentectomy; SL, sublobar; STEPS, Surgical Treatment of Elderly Patients.

while this could be due to confounders, the similar HRs for LCSS largely eliminates greater comorbidities among sublobar resection patients as an explanation. Third, statistically significant differences are seen in most studies involving wedge/sublobar resection vs. lobectomy, and in ~1/3rd of studies involving segmentectomy vs. lobectomy or wedge vs. segment resection. There are no clear additional correlations—results do not seem to track with particular sources of confounding, larger studies, stage, time period or data source.

Several studies (Khullar, Eguchi, Razi) (16,52,53) are categorized as providing high confidence that outcomes are attributable to the resection extent. Two of these found better adjusted OS and LCSS after lobectomy. *Figure 3* shows OS of propensity matched cohorts from the Khullar *et al.* study, which involved extensive matching with several additional analyses (size subsets, margin status, facility type, number of nodes assessed intraoperatively) (16).

On the other hand, Razi *et al.* found no difference in OS for the subset of cIA patients in whom unsuspected pN1 or pN2 nodes were found (52). This study involved extensive adjustment for confounders, including details of the node assessment and use of adjuvant chemotherapy (which

was associated with better OS) (52). Possible reasons for the similar outcomes include that there is no inherent difference between segmentectomy and lobectomy, that any impact of resection extent is overshadowed by that of node involvement, or that a benefit to lobectomy stems from more accurate node assessment and adjuvant chemotherapy (despite being adjusted for). The latter hypothesis is supported by some studies (i.e., similar outcomes with sublobar resection vs. lobectomy when a similar nodal assessment was performed) (61,67,68). However, among adjusted studies overall there is no consistent correlation between long-term outcome differences and adjustment for either adjuvant therapy or extent of node assessment.

Many authors have reported systematic reviews and meta-analyses of non-randomized studies comparing lesser resection to lobectomy (69-75). However, no degree of systematic search rigor or meta-analytic proficiency in amalgamating reported results can overcome residual confounding in the source data. In fact, by combining studies the meta-analytic process obscures the weaknesses of each study. Thus, because of unaccounted (and obscured) confounders, drawing conclusions from meta-analyses of

First author, jow (reference)Study characteristics (reference)Tudy characteristics (reference)Tudy characteristics (reference)Tudy characteristics (reference)Tudy characteristicsTudy characteristicsCaracteristicsSetting characteristicsSetting characteristicsSetting characteristicsSetting characteristicsSetting characteristicsSetting characteristicsSetting characteristicsCaracteristicsSetting characteristicsSetting characteristicsSetting characteristicsSetting characteristicsSetting characteristicsSetting characteristicsSetting characteristi	y vs. lobect						D					000		L \0 -	000
Vis Image $Sidge B E G y vs. lobectStudy chnaracteristicsNorbide spanettingurgerytisticalAdjusSeied % 5 yg vs. LobuNNNAdjustoed % 5 yreg vs. LokeBe$	y vs. lobect	Study ch	naracteristi	cs	Norbid	e span etting urgery	tistical			Adjus Se	ied % 5 y g vs. Lobu	N N N	Adjusto	ed % 5 yr eg vs. Lok	e Be
03-11 418 ^b C A1/2 M M/PM 14/4 H 59 71 14/5 · </th <th>y vs. lobect</th> <th></th> <th>c</th> <th>Stage ^a</th> <th>10D</th> <th>miT e D e D</th> <th>Sta</th> <th></th> <th></th> <th></th> <th></th> <th>HR</th> <th>Seg</th> <th>Lobe</th> <th>HH</th>	y vs. lobect		c	Stage ^a	10D	miT e D e D	Sta					HR	Seg	Lobe	HH
(0:1) (1:1) <th< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></th<>															
0H-13 222 ^b CM1 0H 1H M 71 80 1.11 833 855 0H-13 82 ^b CM2 P PM 11 MM 71 78 1.34 835 855 0H-13 84 ^b CM3 P M 11 M 56. 66 1.72 67 25 75 <			41	clA1,2			MV, PM	14/4	т	59	71	1.45	1	ı	ı
0t-13 222 C/A2 A PM 11 M 57 78 1.34 83 85 0t-13 42 ^b C/A3 A C A C A C A <td< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td>Md</td><td>÷</td><td>Σ</td><td>74</td><td>80</td><td>1.1</td><td>83</td><td>06</td><td>1.32</td></td<>							Md	÷	Σ	74	80	1.1	83	06	1.32
(0-13 442 ⁰ (0,3) (0,1) </td <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>PM</td> <td>Ŧ</td> <td>Σ</td> <td>71</td> <td>78</td> <td>1.34</td> <td>83</td> <td>85</td> <td>1.06</td>							PM	Ŧ	Σ	71	78	1.34	83	85	1.06
02-15 14,286 CM1,2 M <							PM	÷	Σ	50	66	1.72	67	82	1.66
0+15 5,474 CH12 0 0 M 7 7 7 0 0 8 8 8 8 8 8 8 8 8 8 8 8 8 8 1 7 7 1 1 0 9 1 7 7 1 1 9 1 9 1 9 1 9 1 <th< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td>MV, PM</td><td>20/3</td><td>Σ</td><td>65</td><td>68</td><td>1.04</td><td>1</td><td>I</td><td>ı</td></th<>							MV, PM	20/3	Σ	65	68	1.04	1	I	ı
96-06 174 C(A1,2 0 </td <td></td> <td></td> <td></td> <td>cIA1,2</td> <td></td> <td></td> <td>MV, PA, PM</td> <td>8/5</td> <td>Σ</td> <td>76</td> <td>76</td> <td>0.95</td> <td>83</td> <td>83</td> <td>1.02</td>				cIA1,2			MV, PA, PM	8/5	Σ	76	76	0.95	83	83	1.02
0+12 1,637 C/1,12 C 0 0 1 1 7 7 7 1 0 84 66 00-14 1,618° C/1,12 0 0 0 7				cIA1,2			Md	6	_	84	85	1.8	I	ı	ı
00-14 1,618° C A1,2 N M				clA1,2			PM	8/4	_	4	74	1.09	84	86	1.12
05-06 3.500 ClA1,2 MV T L [78] 40 [86] 40 0.83 - 96-04 3.327 ClA1,2 P MV T L 62 ^d T1 ^d 1.04 - - - 96-04 3.327 ClA1,2 P MV T L 62 ^d T1 ^d 1.04 -							MV, PM, IW	11/1	_	74 ^d	76 ^d	1.12	70	74	1.12
96-04 3.327 $(A1,2)$							MV	7	_	[78] ^{d,e}	[86] ^{d,e}	0.83	I		•
03-11 214 $(CA1,2)$ $(CA,2)$				clA1,2			MV	7	_	62 ^d	71 d	1.04	I	I	•
03-13 2.292 ^b clA A MV,PM 6'1 L 66 62 1.08 74 75 03-16 180 ^b clA3 A PA,PM 18 L 58 ^d 61 ^d 1.23 83 88 03-16 180 ^b clA3 A A PM 18 L 58 ^d 61 ^d 1.23 83 88 04-15 1.684 cl-IIA A A A A A A A A A A B A A A B A A B A B A B A B A B A B A B A B A B B A B B A B A B A B A B A B A A B A A				cIA1,2			A	7/1	_	75 ^{d,g}	84 ^d	1.22	1	I	•
03-16 180 ^b cla3 A PA, PM 18 L 58 ^d 61 ^d 1.23 83 88 624 ^b cl-llA A PM 12 L 54 ^{dg} 60^d 1.17 - - 04-15 1,684 clAl A A B VL 76^d 80^d 1.17 - - - 04-15 1,684 clAl A A A B							MV, PM	6/1	_	99	62	1.08	74	75	1.04
624 ^b Cl-IIA A PM 12 L 54 ^{dd} 60 ^d 1.17 - 04-15 1,884 CIA1 A MV 5 VL 76 ^d 80 ^d 1.05 - - 00-12 1,789 CIA1 A MV 6 VL 77 ^d 1.05 81 ^d 87 ^d 00-12 10,500 CIA2 A MV 6 VL 77 ^d 73 ^d 1.22 82 ^d 84 ^d 00-13 46 ^b CIA1,2 A MV 89 VL 67 ^d 73 ^d 1.22 82 ^d 84 ^d 07-13 46 ^b CIA1,2 MV 87 VL 79 ^d 73 ^d 1.27 72 ^d 84 ^d 87 ^d 96-07 14473 Cl-IIA MV 87 ^d VL 79 ^d 1.37 72 ^d 80 ^d 74 ^d 05-13 144 ^f ClIA MV				cIA3			PA, PM	18	_	58 ^d	61 ^d	1.23	83	88	7
$0-15$ $1,684$ $C A1$ A MV 5 VL 76^{d} 80^{d} 1.05 $ 0-12$ $1,789$ $C A1$ A A A A B A B A B MV B VL 78^{d} 1.22 8^{d} 8^{d} $00-12$ $10,500$ $C A2$ P MV B VL 67^{d} 78^{d} 1.27 82^{d} 84^{d} $00-12$ $10,500$ $C A2$ P MV B VL 67^{d} 78^{d} 1.27 82^{d} 84^{d} $00-13$ 462^{b} $C A A$ MV B VL 78^{d} 72^{d} 80^{d} 72^{d} 80^{d} 72^{d} 80^{d} 72^{d} 80^{d} 74^{d} 72^{d} 80^{d} 74^{d} 74^{d} 74^{d} 74^{d} 74^{d} 74^{d} 74^{d} 74^{d} 7		- ت	624 ^b				Μd	12	_	54 ^{d,g}	و0 م	1.17	I	I	
00-12 1.789 C A1 0 MV 6 VL 71 ^d 78 ^d 1.39 81 ^d 87 ^d 00-12 10,500 C A2 0 MV 8 VL 67 ^d 73 ^d 1.22 82 ^d 84 ^d 98-07 5,118 C A1,2 MV 9 VL 58 ^d 70 ^d 1.37 72 ^d 84 ^d 07-13 46 ^z c-IIA MV 9 VL 58 ^d 70 ^d 1.37 72 ^d 84 ^d 98-07 14,473 cl-IIA MV 9 VL 79 78 1.65 - - - - 98-07 14,473 cl-IIA MV 9 VL 79 78 1.65 ^d 74 ^d - - <td< td=""><td></td><td></td><td>1,6</td><td>cIA1</td><td></td><td></td><td>M</td><td>ъ</td><td>٧L</td><td>76 ^d</td><td>80 ^d</td><td>1.05</td><td>1</td><td>I</td><td>ı</td></td<>			1,6	cIA1			M	ъ	٧L	76 ^d	80 ^d	1.05	1	I	ı
00-12 10,500 C/A2 A MV 8 VL 67 ^d 73 ^d 1.22 82 ^d 84 ^d 98-07 5,118 C/A1,2 P MV 9 VL 58 ^d 70 ^d 1.37 72 ^d 80 ^d 07-13 46 ² C/-IIA MV 9 VL 79 ^d 73 ^d 1.65 5 ^d 80 ^d 98-07 14,473 CI-IIA MV 9 VL 79 ^d 78 ^d 165 ^d 63 ^d 74 ^d 98-07 14,473 CI-IIA MV 9 VL 79 ^d 78 ^d 165 ^d 5 ^d 74 ^d 98-07 14,473 CI-IIA MV 9 VL 79 ^d 29 ^d 74 ^d 74 ^d 06-13 188 ^b CI,II P P VL 94 ^d 96 ^d 74 ^d 74 ^d 18. 18. ^d MV P VL 96 ^d 96 ^d 74 ^d 74 ^d <td></td> <td></td> <td>1,7</td> <td>cIA1</td> <td></td> <td></td> <td>MV</td> <td>9</td> <td>٨L</td> <td>71 ^d</td> <td>78 ^d</td> <td>1.39</td> <td>81 ^d</td> <td></td> <td>1.64</td>			1,7	cIA1			MV	9	٨L	71 ^d	78 ^d	1.39	81 ^d		1.64
98-07 5,118 ClA1,2 MV 9 VL 58 ^d 70 ^d 1.37 72 ^d 80 ^d 07-13 462 ^b Cl-IIA P PM 5 VL 79 78 1.65 - - - 98-07 14,473 Cl-IIA PM 5 VL 79 78 1.65 - - - - 98-07 14,473 Cl-IIA PM 9 VL 79 ^d 80 ^d 74 ^d - -							M	∞	٨L	97 ^d	73 ^d	1.22	82 ^d	84 ^d	1.13
07-13 462 ^b Cl-IIA PM 5 VL 79 78 1.65 - - 98-07 14,473 Cl-IIA MV MV 9 VL 50 ^d 62 ^d 1.37 63 ^d 74 ^d 05-13 188 ^b cl-IIA MV PM 7 VL 50 ^d 62 ^d 1.37 63 ^d 74 ^d 05-13 188 ^b cl.II MV PM 7 VL [94] ^{6,9} [96] ⁶ - - - - 05-15 454 ^b ClA, pN1 ^h MV PAPM 19 VH [42] ^h [44] ^h 0.92 - - - 05-15 430 ^b ClA, pN2 ⁱ MV PAPM 19 VH [42] ^h 1092 - <t< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td>MV</td><td>6</td><td>٧L</td><td>58 ^d</td><td>20 d</td><td>1.37</td><td>72 ^d</td><td>80 ^d</td><td>1.37</td></t<>							MV	6	٧L	58 ^d	20 d	1.37	72 ^d	80 ^d	1.37
98-07 14,473 Cl-IIA MV 9 VL 50 ^d 62 ^d 1.37 63 ^d 74 ^d 05-13 188 ^b cl,II MV PM 7 VL [94] *9 [96] * - <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>ΡM</td> <td>5</td> <td>٧L</td> <td>79</td> <td>78</td> <td>1.65</td> <td>I</td> <td>I</td> <td>ı</td>							ΡM	5	٧L	79	78	1.65	I	I	ı
05-13 188 ^b CI,II E PM 7 VL [94] ^{6,9} - - </td <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>M</td> <td>6</td> <td>٧L</td> <td>50 ^d</td> <td>62 ^d</td> <td>1.37</td> <td>63 ^d</td> <td>74 ^d</td> <td>1.37</td>							M	6	٧L	50 ^d	62 ^d	1.37	63 ^d	74 ^d	1.37
05-15 454 ^b CIA, pN1 ^b MV PA PM 19 VH [42] ^h [44] ^h 0.92 - - 05-15 430 ^b CIA, pN2 ⁱ MV PA PM 19 VH [42] ⁱ [37] ⁱ 1.09 - -							ΡM	7	VL	[94] ^{e,g}	。[96]	1	I	I	I
NCDB 05-15 454 ^b CIA, pN1 ^b MV PA PM 19 VH [42] ^h [44] ^h 0.92 - - - NCDB 05-15 430 ^b cIA, pN2 ⁱ MV PA PM 19 VH [42] ⁱ [37] ⁱ 1.09 - - -	Segment vs. lobectomy, pN po	sitive													
NCDB 05-15 430 ^b CIA, pN2 ¹ MV PA PM 19 VH [42] ¹ [37] ¹							MV PA PM	19	ΗΛ	[42] ^h	[44] ^h	0.92	I	ı	ı
							MV PA PM	19	ΗΛ	[42]	[37]	1.09	I	I	1

Journal of Thoracic Disease, Vol 14, No 6 June 2022

 Table 2 Long-term outcomes in generally healthy patients: sublobar or wedge resection vs. lobectomy

 Ordered by resection extent, deeree of confidence that results reflect the effect of the treatment, stage

					Adjustme	ent for co	Adjustment for confounding								
First author, year (reference)	Ó	tudy char	Study characteristics	(0	nogr F tage e span tting	ţnwoı. nıdeu)	istical hods	it for/ sets	The street and the st	Adjust SL/	Adjusted % 5 yr OS SL/W vs. Lobe	rr OS be	Adjust SL	Adjusted % 5 yr LCSS SL/W vs. Lobe	LCSS
	Source	Yrs	c	Stage ^a	AoD e iH miT	Q si				SL/W	Lobe	HR	SL/W	Lobe	HH
Sublobar resection vs.	lobectomy														
Eguchi 2019 (53)	US ×1	95-14	698 ^b	с			Μd	19/4	т	78	82	~	91	94	1.95
Yu 2020 (54)	SEER	04-13	462 ^b	cIA1,2 ¹			MV, PA, PM	15/3	_	53	68	1.38	63	79	1.45
Eguchi 2017 (55)	US ×1	00-11	2,186	cl-IIA			M	12/1	_	67 ^d	78 ^d	1.74	86 ^d	91 d	2.06
Liang 2019 (56)	SEER	04-14	22,914	G			M	8	٨L	I	ı	ı	71	82	1.57
Wedge resection vs. Ic	lobectomy														
Dolan 2021 (57)	US ×1	10-16	1,086	cl			MV, PA	25/2	ΗΛ	83	86	1.23	1	ı	
Boyer ^k 2017 (58)	Ν	01-10	3,196 ^b	cl-IIA			MV, PA	8/6	т	44	52	1.22	1	ı	
Khullar 2015 (16)	NCDB	03-11	418 ^b	cIA1,2			MV, PM	14/4	Σ	55	71	1.7	1	ı	1
Cao 2018 (36)	SEER	04-13	1,028 ^b	cIA1			M	÷	_	74	80	1.2	84	89	1.3
Cao 2018 (36)	SEER	04-13	3,362 ^b	cIA2			M	Ē	_	63	75	1.58	27	85	1.66
Cao 2018 (36)	SEER	04-13	1,298 ^b	cIA3			Mq	11	_	48	65	1.63	65	73	1.46
Yendamuri 2013 (42)	SEER	05-08	3,509	cIA1,2			M	7	_	[82] ^{d,e}	[86] ^{d,e}	1.09	1	I	I
Yendamuri 2013 (42)	SEER	98-04	3327	cIA1,2			M	7	_	53 ^d	71 d	1.19	1	I	I
Speicher ^k 2016 (59)	NCDB	03-06	11,990	cIA			M	6/2	_	51 ^d	و و م	1.52	1	I	ı
Subramanian ^k 18 (60)	NCDB	06-07	325 ^b	cIA			ΡM	16	_	56 ^d	62 ^d	1.18	1	I	I
Fan 2020 (47)	SEER	04-15	2,360	cIA1			M	5	٧L	71 ^d	80 d	1.36	1	I	I
Dai 2016 (48)	SEER	00-12	2,450	cIA1			M	9	٧L	68 ^d	78 ^d	1.45	83 ^d	87 ^d	1.45
Dai 2016 (48)	SEER	00-12	12,386	cIA2			M	8	٧L	62 ^d	73 ^d	1.64	73 ^d	84 ^d	1.68
Cox ^{k,m} 2017 (61)	NCDB	03-06	1,191	cl-IIA			M	4/1	٧L	68 ^d	71 d	1.23	1	I	ı
Dziedzic 2017 (50)	Polish Reg	07-13	462 ^b	cl-IIA			ΡM	5	٧L	54	78	2.5	1	I	ı
Nakamura ^f 2011 (62)	Japan ×1	<i>2-</i> 00	373	cl-IIA			MV	4	٧L	55 ^d	82 ^d	4.3	I	ı	ı
Inclusion criteria: studies with multivariable or propensity adjustment of sublobar or wedge resection vs. lobectomy, 2000-21, with >50 pts per arm in generally healthy	lies with mu	ltivariable	er prope	ensity adjus	tment of subloba	r or wed	ge resectior	n vs. lob	ectomy, 1	2000–21,	with >5(0 pts pe	r arm in g	generally	nealthy
patients with generally solid tumors; excluding studies that accrued most patients before 2000. The HR reference is lobectomy, i.e., HR >1 reflects worse outcome compared	solid tumors	s; excludi	ng studie:	s that accru	ed most patients	before 20 -hadina	00. The HR	referenc	e is lobe	ctomy, i.e	., HR >1	reflects	worse ou	tcome col	npared
wur tobectorny. Bota nigningnis beuer outcorne (>∠-point anierence); Lignt green snaaing nigniignus stausucany signiincant anierence (lignter snade = univariable; darker = multivariable). For abbreviations, footnotes, explanation of adjustment for confounding see legend for <i>Table</i> 3.	rigriignis be	otnotes.	orne (>z-f explanatio	point annere in of adiustr	>z-point anierence); Light green shaaing nighiights staust ation of adjiistment for conforinding see legend for <i>Table</i> 3	snauing ing see le	nigniignis s dend for <i>Tal</i>	lausucai 5/e 3.	y signilio	cant oner	ence (iig	mer sna	de = univ	ariable; d	arker =
						, , , , , , , , , , , , , , , , , , ,	5. 12. 10.								

	stage
segmentectomy	ct of the treatment, st
1 <i>VS</i> .	ect
ctior	e eff
rese	lect the effec
edge	s reflec
ts: wed	sults
tien	at re
thy pa	e thi
ealth	denc
lly h	onfie
neral	ofc
n gei	gree
ies i	t, de
itcomes in ger	xten
n o n	one
g-terr	secti
Ë	ov re
3 Lo	ed b
Fable	Ordered
C	

			:		Ac	Adjustment for confounding	or confe	ounding								
First author, year	U)	study char	Study characteristics		bio	б	IOL			d RE	Adjust V	Adjusted % 5 yr OS W vs. Seg	r OS	Adjuste	Adjusted % 5 yr LCSS W vs. Seg	LCSS
(reierence)		:	:	(nogi Norb tage	ettin ettin	աու ,	tistic bodi	iot įb etece							
	Source	Yrs	z	Stage ^ª	10D	Q S	νвЯ				Μ	Seg	HH	×	Seg	HR
Wedge resection vs. segmentectomy	egmentecton	ny														
Smith ⁿ 2013 (63)	SEER	98-06	3,525 ⁿ	cIA1,2			PA	PA, PQ, PM	7/2	Σ	1		1.19	I	ı	1.22
Smith ⁿ 2013 (63)	SEER	98-06	3,525	clA			PA	PA, PQ, PM	7/2	Σ	I	I	1.23	I	I	1.32
Koike 2013 (64)	Japan ×1	60-86	328	cIA				MV	15	Σ	ı	ı	1	68 ^d	91 d	3.18
Cao 2018 (36)	SEER	04-13	252 ^b	cIA1				M	÷	_	76	74	1.05	83	91	.75
Cao 2018 (36)	SEER	04-13	852 ^b	cIA2				M	11	_	64	72	1.34	75	85	1.65
Cao 2018 (36)	SEER	04-13	440 ^b	cIA3				M	11	_	48	53	1.17	62	69	1.25
Zhang ° 2016 (65)	SEER	98-12	3,391	cIA				PA	8/2	_	ı	ı	1.15	I	ı	1.09
Zhang ^p 2016 (65)	SEER	98-12	1,949	cIA				PA	8/2	_	ı	ı	F	ı	ı	.92
Fan 2020 (47)	SEER	04-15	1,026	cIA1				MV	5	٨L	71 ^d	76 ^d	1.42	I	1	ı
Dai 2016 (48)	SEER	00-12	981	cIA1				MV	9	٧L	68 ^d	71 ^d	1.08	83 ^d	81 ^d	.93
Dai 2016 (48)	SEER	00-12	3,104	cIA2				MV	ø	٧L	62 ^d	67 ^d	1.36	73 ^d	82 d	1.42
Zhao 2019 (66)	SEER	04-15	1,372 ^b	cIA			~	MV, PM	10/3	٧L	39	68	1.29	77	78	I
Dziedzic 2017 (50)	Polish Reg	07-13	462 ^b	cl-IIA				PM	5	VL	54	79	1.49	I	I	I
Inclusion criteria: studies with multivariable or propensity adjustment of wedge resection vs. segmentectomy, 2000-21, with >50 pts per arm in generally healthy patients with	ies with multi	ivariable c	or propens	ity adjustm	ent of wedg	ge resectior	J VS. Se	gmentect	omy, 200	0–21, wi	th >50 pt	ts per arn	n in gene	rally hea	Ithy patien	ts with

generally solid tumors; excluding studies that accrued most patients before 2000. The HR reference is segmentectomy, i.e., HR >1 reflects worse outcome compared with segmentectomy. Bold highlights better outcome (>2-point difference); Light green shading highlights statistically significant difference (lighter shade = univariable; darker = multivariable)

_egend (*Tables 1-3*):

30-50% were "lobe-like" segments (lingula-sparing Left Upper Lobectomy, lingulectomy or basilar quadri-segmentectomy); ", cN0 but pN1 (OS in brackets because not 8th edition stage classification (reported stage is translated into current 8th edition nomenclature for the sake of uniformity and contemporary application); ^b, propensity comparable to unselected cN0 cohorts; cn0 but pN2 (OS in brackets because not comparable to unselected cn0 cohorts; all with visceral pleural invasion (technically stage IB but ≤2 cm); ^k, predominantly wedge (≥80%); ^l, ACS special study (involving enhanced chart abstraction of clinical factors); ^m, lepidic adenocarcinoma; ⁿ, for entire matched pairs (total); ^c, all solid tumors (GGN excluded); ^d, unadjusted results; ^d, 3-year survival (in brackets because not comparable to 5-year OS); ¹, All resected by VATS; study, not this specific cohort; °, adenocarcinoma; P, squamous carcinoma.

HR, hazard ratio; LCSS, lung cancer specific survival; Lobe, lobectomy; NCDB, US national cancer database; NS, not statistically significant; OS, overall survival; Reg, registry; SEER, Surveillance, Epidemiology, and End Results database; Seg, segmentectomy; SL, sublobar resection (segmentectomy or wedge); STS-MC, Society of thoracic Surgeons Database, linked to Medicare; VATS, video-assisted thoracic surgery; W, wedge; Yrs, years (of patient accrual).

span, adjustment for changes during the study period or differential use of the interventions; Q settings, discrepancy in the facilities or settings performing the interventions; Q treatmt, quality of the treatment (e.g., margin distance, adjuvant therapy); Fav tumor, selection of less aggressive tumors for an intervention; Statistical methods, methods used to adjust for confounding; Subset, additional Adjustment for Confounding: Demogr F, demographic factors (age, sex, socioeconomic); CoMorbid, comorbidities; Hi stage, occult stage inaccuracy due to differences in extent of assessment; Time subset or sensitivity analyses, # adj for, number of factors adjusted for, Conf RE that effect, Confidence that results reflect the effect of the treatment vs. confounding factors. MV, Multivariable model (e.g.,

	Clearly confounded	VL-very low confidence
	High concern	VL-very lo
re quintiles.	Moderate concern	L-low
ty matching; PQ, analysis of propensity score quintiles.	Limited concern	M-moderate
atching; PQ, analys	Neutral (likely little effect)	H-high
PM, propensit	Addressed	VH-very high
ssion); PA, propensity score adjustment;	Categories of confounding	Confidence RE treatment effect
Cox regression	Color	code:

2366

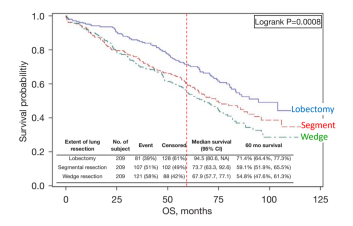


Figure 3 Propensity-matched comparison of wedge resection, segmentectomy and lobectomy.

Comparison of resection extent in the National Cancer Database of cIA1,2 NSCLC [2003–6]. This study matched for 14 prognostic factors and performed multiple sensitivity tests; it is assessed to have a low level of residual confounding. Reproduced with permission from Khullar *et al.* (16). OS, overall survival.

non-randomized studies is problematic.

Recurrence

Recurrence is a concern, especially because the LCSG821 RCT found a higher local recurrence rate after sublobar resection (8,31). However, assessment of this outcome is impacted by multiple factors (e.g., competing causes of death, length of follow-up, staging accuracy, tumor biology). The cleanest measure is FFR (or cumulativeincidence-of-recurrence). Recurrence-free or disease-free survival (RFS/DFS) is muddy because it mingles recurrence with competing causes of death. Simple comparison of the number (or type) of observed recurrences in cohorts is frequently reported but hard to interpret (no accounting for confounding factors or follow-up duration).

Few adjusted NRCs report recurrence by resection extent (*Table 4*) (39,43,45,46,53,57,60,64,76-80). The available evidence is unclear whether lesser resection increases recurrence risk. The confidence that confounders are accounted for is low. Variability in the incidence of recurrence is only partially potentially explained by tumor stage or follow-up duration. Most studies found a nonsignificant trend towards a higher recurrence rate after sublobar resection, rarely the opposite trend. Rates of locoregional recurrence are generally low (the outcome

Detterbeck et al. Evidence for resection extent in healthy patients

most likely affected by resection extent).

Pulmonary function tests (PFTs)

The impact of resection on PFTs serves as a surrogate for functional capacity (which hasn't been studied). Segmentectomy doesn't confer a meaningful benefit over lobectomy in healthy patients; studies reporting FEV1 ≥ 6 months postoperatively are shown in *Table 5* (changes in diffusion capacity are seldom reported) (8,29,35,51,81-96) (it takes ~6 months following surgery for PFTs to reach a plateau; less after VATS resection) (95,97-99).

Lobectomy causes a ~14% long-term decrease in FEV1. Segmentectomy results in an FEV1 decrease of ~12% in studies involving many multi-segment resections (e.g., left upper tri-segmentectomy) and a decrease of ~5% in studies involving primarily single segment resections. Such decreases are not in a clinically relevant range for healthy patients. Indeed, exercise capacity is reported unchanged despite the FEV1 decrease (83,91). Available data shows an FEV1 decrease of 2–8% after wedge resection (89,95,100,101). The long-term impact of resection on FEV1 does not correlate with the time period or the approach (VATS/open).

Long-term QOL

In *Table 6* (102-117) and *Table 7* (11,24,118-130) postoperative QOL results are depicted reflecting no change, or small, moderate or large changes *vs.* baseline by generally accepted thresholds for clinically meaningful differences (128,131-136). *Table 6* is mostly yellow (i.e., no change); these studies used the SF-36 tool (why this tool appears less sensitive is unclear; little change remains when using lower proposed thresholds for clinically meaningful differences). In *Table 6* and *Table 7*, there is diminishing QOL impairment towards the right (i.e., increasing interval from surgery) and increasing impairment moving downward. The vertical gradient reflects increased VATS near the top and more extensive resections (e.g., pneumonectomy) towards the bottom (also generally older studies).

What conclusions can be drawn? The SF-36 tool seems less useful. VATS is associated with less QOL impairment vs. baseline, and this has mostly resolved by 6 months (except dyspnea). Whether sublobar resection has an impact is less clear—studies are limited and confounded by the use of VATS. Open lobectomy is associated with long-term QOL decreases in many domains. Older studies tend to show larger and more frequent QOL impairment, but often

1 st author, year (reference)		Stud	y charao	Study characteristics		Afid RE It effect	ration of (mo)	Unmatched overall recurrence %	tched rall nce %	Unmatched locoregional recurrence %	ched jional nce %	Adjusted RFS/DFS Seg/W <i>vs</i> . Lobe	sted DFS s. Lobe	Adjusted FFR Seg/W vs. Lobe	sted R s. Lobe
	Source	Yrs	ч	Lobe vs.:	Stage ^a			Seg/W	Lobe	Seg/W	Lobe	HR	Ъ	HR	Р
Lesser resection vs. lobectomy	ctomy														
Dolan 2021 (57)	US×1	10-16	1,086	>	cl	ΗΛ	51	24 ^b	11 ^b	13 ^b	2 p	1.4	SN	I	
Eguchi 2019 (53)	US×1	95-14	° 869	SL	С	т	ı	18 ^b	а 6	10	0	ı	ı	2.33	<.001
Koike ^d 2016 (39)	Japan ×1	98-09	174	Seg	cIA1,2	_	78	23 ^b	20 ^b	10 ^b	°	1.5	SN	I	ı
Chan 2021 (45)	US×1	03-16	180 °	Seg	cIA3	_	60	24	23	12	6	1.23	SN	1.05	NS
Landreneau 2014 (46)	US×1	ı	624 °	Seg [°]	cl-IIA	_	65	20	17	9	5	ı	I	1.11	NS
Subramanian 2018 (60)	NCDB ¹	06-07	325 °	M ^g	cIA	_	>60	1	I	ı	1	1	I	1.39	<.05
Huang 2020 (76)	China ×1	06-16	238 °	SL	pIA ^h	_	65	1	I	I	I	.85	SN	I	I
Yamashita 2012 (43)	Japan ×1	03-11	214	Seg°	cIA1,2	٨L	30	œ	9	4	ю	1.12	NS	I	I
Kamigaichi 2020 (77)	Japan ×3	10-16	230 °	Seg [°]	cIA1,2	٨L	37	ß	11	5	7	$\overline{\nabla}$	SN	7	NS
El-Sherif 2006 (78)	US×1	90-03	784 °	SL	cI-IIA	٨L	31	29	28	7 J	4	1.2	NS	I	I
Wedge resection vs. segmentectomy	mentectomy							M	Seg	M	Seg	W vs.	Seg	W vs.	Seg
Tsutani ^{k,I} 2021 (79)	Japan ×3	10-15	457	Seg vs. W	cIA	н	48	13 ^b	7 ^b	•	I	ı	ı	2.13	.02
Altorki ^k 2016 (80)	US×1	00-14	289	Seg vs. W	cIA	Σ	34	19	20	÷	ര	1.05	SN	I	I
Koike 2013 (64)	Japan ×1	98-09	328	Seg vs. W	cIA	Δ	58	I	I	34	6	I	ı	5.79	<.001
Inclusion criteria: studies reporting RFS, DFS or	s reporting F	RS, DFS		FFR with multivariable or propensity adjustment of segmentectomy or wedge resection vs. lobectomy, 2000-21, with ≥50	iable or pro	pensity a	djustme	nt of segn	nentecton	ny or wedg	je resecti	on vs. lob	ectomy, 2	000-21, \	with ≥50

 Table 4 Recurrence outcomes in generally healthy patients

 Ordered by resection extent, degree of confidence that results reflect the effect of the treatm

patients per arm in generally healthy patients with generally solid tumors. The HR reference is lobectomy, i.e., HR >1 reflects worse outcome compared with lobectomy. Bold nighlights better outcome (>2-point difference); Light green shading highlights statistically significant differences (lighter shade = univariable; darker = multivariable); Red font 8th edition stage classification (reported stage is translated into current 8th edition nomenclature for the sake of uniformity and contemporary application); ^b, matched cohort; ^c nighlights accrual occurring primarily before 2000.

propensity matched pairs (total); ^d, all solid tumors (GGN excluded); ^o, 30–50% were "lobe-like" segments (lingula-sparing left upper lobectomy, lingulectomy or basilar quadri-~25% predominantly ground glass but excluded AIS & MIA; | solid tumor size, CTR ≥0.8, PET SUV ≥2.5; | local only (adjacent lung parenchyma); ^k, excluded AIS, MIA; | ~50% , solid tumor size, segmentectomy); ¹, American College of Surgeons special study (involving enhanced chart abstraction of clinical factors); ^a, predominantly wedge (≥80%); ¹ nad minor GG component.

AS, adenocarcinoma in situ; Conf RE tmt effect, Confidence that results reflect the effect of the treatment (lobectomy or SL resection) vs. confounding factors; DFS, disease ree survival; FFR, freedom from recurrence (only recurrence counts as an event); f/u, follow up duration (months); HR, hazard ratio; L, low confidence; Lobe, lobectomy; M, moderate confidence; MIA, minimally invasive adenocarcinoma; NCDB, US national cancer database; NS, not statistically significant; RFS, recurrence free survival; Seg, segmentectomy; SL, sublobar resection (segmentectomy or wedge); W, wedge; VH, very high confidence; VL, very low confidence; Yrs, years (of patient accrual).

Journal of Thoracic Disease, Vol 14, No 6 June 2022

2368

1 st author, year (reference)	Years	N Lobe/Seg	Open/VATS	Interval to PFT		erence in FEV		Comments
(reierence)		Lobe/Seg		(mo)	Seg	Lobe	Р	
Frequent ^a multi-segmer	ntectomy							
Yoshikawa 2002 (81)	1992-94	55	Open	12	-13%	-	-	
Takizawa 1999 (82)	1993-96	40/40	Open	12	-7%	-14%	<0.05	
Harada 2005 (83)	-	45/38	Open	6	-12%	–18% ^b	<0.05	
Kashiwabara 2009 (84)	2000-06	20/30	Open	6	-14%	-13%	NS	Preop FEV1 <70%
Kashiwabara 2009 (84)	2000-06	27/41	Open	6	–13%	-19%	<0.05	Preop FEV1 >70%
Yoshimoto 2009 (85)	2005-07	-/56	Open	12	-12%	-	-	
Saito 2014 (86)	2006-12	126/52	Open	6	-10%	–19% ^b	NS	
Nomori 2016 (87)	2013-15	13/20	Open	7	-10%	-17%	<0.05	≥2 segments
Hwang 2015 (51)	2005-13	94/94	VATS	?	-9%	–11% [⊳]	NS	
Handa 2019 (29)	2007-17	-/50	VATS	12	-11%	-	-	2 segments
Suzuki 2017 (88)	2009-12	33/37	VATS	>6	-12%	–11% [⊳]	NS	
Saji 2022 (35)	2009-14	526/528	VATS	12	-9%	-12%	<.0001	
Gu 2018 (89)	2011-14	75/34	VATS	6	-18%	-21%	NS	
Tane 2020 (90)	2012-17	88/35	VATS	6	-12%	-18%	-	Left upper division
Subset		-			-12%	-16%	2 	
Few multi-segment rese	ctions					*	•	
Ginsberg 1995 (8)	1982-88	67/71	Open	6	-2%	-9%	<0.05	1/3 rd wedge
Keenan 2004 (91)	1996-01	147/54	Open	12	-5%	–11% ^b	-	
Nomori 2012 (92)	2005-09	-/96	Open	6	-10%	-	-	
Nomori 2016 (87)	2013-15	13/83	Open	7	-2%	-17%	<0.05	1 segment
Nomori 2018 (93)	2013-16	103/103	Open	7	-5%	-13%	<0.05	
Macke 2015 (94)	2002-10	82/77	VATS °	>6	-4%	-8%	<0.05	1–2 vs. 3–5 segments
Kobayashi 2017 (95)	2001-9	228/118	VATS ^d	12	-7%	–10% ^b	-	
Handa 2019 (29)	2007-17	-/88	VATS	12	-10%	-	-	1 segment
Helminen 2020 (96)	2007-19	48/50	VATS	~9	+1%	-8%	<0.001	
Tane 2020 (90)	2012-17	88/23	VATS	6	-5%	-18%	-	1 segment
Subset					-5%	-12%	-	
Average					-9%	-14%		

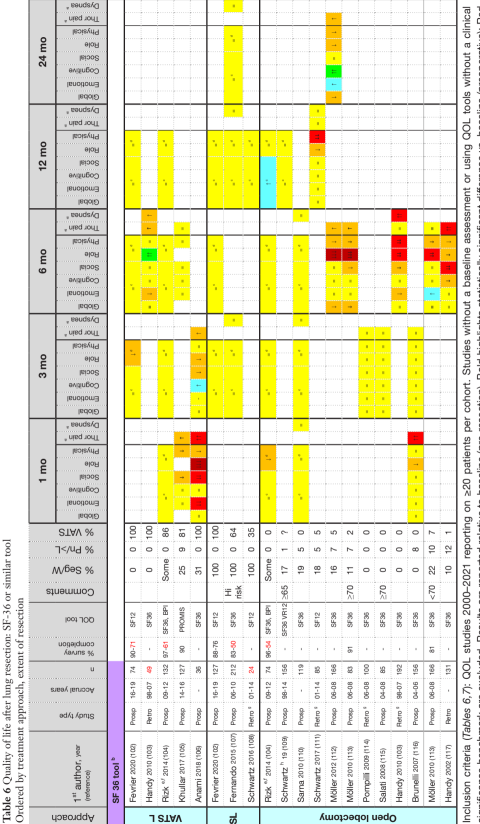
Table 5 Change in lung function following segmentectomy or lobectomy

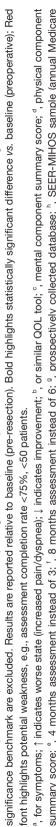
 Ordered by single/multi-segmentectomy, VATS/open approach, years of accrual

Inclusion criteria: studies involving sublobar resection reporting a change in pulmonary function tests, published 1995–2021, ≥50 patients total; Red font highlights accrual occurring primarily before 2000. Light yellow shading highlights major focus of table.

^a, including >30% "lobe-like" segmentectomies (left upper trisegmentectomy, lingulectomy or basilar multi-segmentectomy; ^b, lobectomy included RML; ^c, mostly VATS; ^d, lobectomies were mostly VATS, segmentectomies mostly open.

FEV1, forced expiratory volume in 1 second; Lobe, lobectomy; mo, months; NS, not statistically significant; PFT, pulmonary function test; Preop, preoperative; Seg, segmentectomy; RML, right middle lobectomy; VATS, video-assisted thoracic surgery.





¹, 4 months assessment instead of 3; ¹, 8 months assessment instead of 6; ⁹, prospectively collected database; ¹, SEER-MIHOS sample (annual Medicare Hi risk, patients deemed unfit to tolerate lobectomy by ACOSOG hi risk criteria; Lobe, lobectomy; Pn/>L, pneumonectomy or extended lobectomy (e.g., bilobectomy, + chest Outcomes Survey conducted in a representative sample); ¹, average of the 2 cohorts; ¹, for total group, not necessarily this subset; ^k, cohort without recurrence. summary score;

vall, sleeve resection); Prosp, prospective; QOL, quality-of-life; RCT, randomized controlled trial; Retro, retrospective; Seg, segmentectomy; SL, sublobar resection; Thor, thoracic; VATS, video-assisted thoracic surgery; W, wedge resection. ⁻or QOL color assessment code see legend for Table 7.

Dyspnea I you bain Physical 0 E Bole 24 Social Cognitive Emotional Global Dyspnea Thor pain Physical 0 E Bole 42 Social Cognitive Emotional Global Dyspnea ^{*} Thor pain Physical 6 mo Bole Social Cognitive Emotional Global Dyspnea Thor pain Physical 3 mo Bole Social . Cognitive п Emotional Global Dyspnea Thor pain Physical 0 E Bole Social -Cognitive ю. Emotional Table (Global 100 100 100 92 85 68 32 43 ₽ 0 For inclusion criteria, abbreviations and footnotes see legend for STAV % 59 23 0 0 N 0 0 0 0 0 0 Table 7 Quality of life after lung resection: EORTC or similar tool 9 100 /v₽ % 0 0 ω 0 0 0 0 0 0 0 0 23 0 0 6 9 9 ω 100 100 W/ge2 % 0 ī ÷ 25 0 0 0 N 0 0 0 0 0 0 0 0 0 0 ı. 270 >70 <70 Seg >70 <70 Comments ≥ Ordered by treatment approach, extent of resection EQ5D C30 LC13 C30, LC13 C30, LC13 C30, LC13 C30, LC13 C30, LC13 C30, LC13 EQ5D C30, LC13 C30, LC13 C30, LC13 EQ5D C30, LC13 C30, LC13 C30, LC13 C30, LC13 15D C30 C30 C30 QOL tool C30, c30, C30, 91-<mark>68</mark> 98-<mark>73</mark> completion % survey 95<mark>-68</mark> 3-68 80-<mark>62</mark> 80-72 86-73 93-85 91-82 22-06 84-<mark>69</mark> 98-<mark>73</mark> 95-80 94-<mark>68</mark> 90-76 90-73 75 10 8 96 166 256 u 120 102 99 74 88 ដ 5 54 66 19 61 131 4 68 53 111 58 8 4 02-04 08-14 03-06 02-04 14-16 14-16 14-15 13-16 13-16 98-04 98-04 08-14 99-05 02-04 98-04 98-04 02-05 00-66 99-05 03-08 Ассгиаl уеага 4 Prosp Retro⁹ Prosp Prosp Retro⁹ Prosp rosp Prosp rosp rosp Prosp Prosp Prosp Prosp rosp RCT Prosp RCT Retro RCT RCT Study type Benedixen 2016 (24) Balduyck 2007 (123) Benedixen 2016 (24) Balduyck 2009 (125) Balduyck 2008 (130) Balduyck 2007 (123) Burfeind 2008 (122) Burfeind 2008 (122) Kenny ^{e,k} 2008 (129) Stamatis 2019 (11) Stamatis 2019(11) Schulte 2009 (126) Pompili 2018 (119) Schulte 2009 (126) Schulte 2010 (127) Schulte 2010 (127) year Nugent 2020 (120) Alberts 2019 (124) llonen 2010 (128) Avery 2020 (121) EORTC tool Xu¹ 2020 (118) author, (apr 1st refe Ч Approach **J STAV** SL Open lobectomy

QOL asse	OL assessment color code:		
111	>20 points* better	2x clinically meaningful improvement	* for normalized QOL scales a 10-point difference is usually accepted as clinically meaningful (C-30, LC-13,
ţţ	10-20 points* better	Clinically meaningful improvement	EGOD, OF-50, FROMIS, OTHER SCARES AURITIED TO COTIESPOND
←	5-<10 points* better	Somewhat better	Mapping of SF38: General health = global; role emotional = emotional; mental health = cognitive; social function-
II	Same (0-<5 points*)	Similar to baseline (i.e., pre-treatment)	ing = social; role physical = role; physical tunctioning = physical; bodily pain = thoracic pain; 15D: Total = diobal: Depression/distress = emotional: mental functioning = coonitive: usual activities = role:
→	5-<10 points* worse	Somewhat worse	Mobility = physical; discomfort = pain; breathing = dyspnea
11	10-20 points* worse	Clinically meaningful worsening	PROMIS: Anxiety/depression/emotional support = emotional; informational support = cognitive; social roles = social: physical function = physical: pain intensity/intenference = pain
111	>20 points* worse	2x clinically meaningful worsening	EQ5D: Health index = global; anxiety/depression = emotional; usual activities = role; mobility = physical;

include larger resections.

The average doesn't necessarily reflect an individual's experience. Another measure is the proportion of patients that have improved, unchanged or worse QOL after surgery. Six months after thoracotomy, one study reported that 30-50% of patients experience meaningfully worse QOL vs. baseline (SF-36 instrument, included 9% pneumonectomy) (137). In another study, long-term QOL after thoracotomy was meaningfully worse in ~10-40% and improved in a similar proportion in various domains of the EORTC C-30 instrument in patients without recurrence (129). These authors reported that long-term symptoms were absent or meaningfully improved in ~60% and worse in ~10-20%-with the exception of dyspnea which was worse in ~40% vs. baseline. A prospective study involving primarily minimally invasive resections found that 20-40% were meaningfully worse and a similar proportion improved at 6 and 12 months in multiple EORTC domains (120). No data is available whether these proportions are influenced by sublobar resection.

Various predictors of worse QOL have been noted, mostly in single studies and measures of physical functioning. Worse long-term QOL has been associated with age (137) smoking (138), adjuvant chemotherapy (137), recurrence (129), higher baseline QOL (139), thoracotomy (vs. VATS) (111) and larger resection (i.e., pneumonectomy or lower ppoFEV1) (137,139). One study noted a nonsignificant trend to less impact on QOL with sublobar resection vs. lobectomy (137); another found physical QOL at ~11 months was unchanged after limited resection but decreased after lobectomy (likely confounded by use of VATS) (108,111). Conversely, variables that don't correlate with QOL changes include gender (112,140), comorbidities, occurrence of postoperative complications, and stage (137). A case-matched study found no association between the presence of COPD and postoperative QOL (114).

Two recent small RCTs deserve mention. A RCT of lobectomy (VATS vs. open) found a transient QOL impairment with return to baseline or higher; the return was faster after VATS (6 vs. 12 weeks) (24). A small RCT of segmentectomy vs. lobectomy found that global QOL returned to baseline by 3 months in both arms (11). Interpretation is difficult, however, because of the study size (n=108) and higher VATS use in the lobectomy arm (11).

Chronic pain

The incidence of chronic pain is reported variably. The impact of sublobar resection is unclear, confounded

by VATS use. No differences were found in one study of 220 patients undergoing either VATS lobectomy, segmentectomy via mini-thoracotomy, or segmentectomy via thoracotomy with rib-spreading [2012–5]. At 1 month ~25% in each group were taking analgesics (of any kind), and by 3 months it was $\leq 5\%$ (28). Moderate to severe pain persisted in 5–10% of patients at 1 year in a RCT of VATS *vs.* open lobectomy but was approximately half as frequent after VATS (24). In *Table 6* and *Table 7*, pain at ≥ 6 months postoperatively is noted frequently after thoracotomy but infrequently after VATS.

Several studies addressing chronic pain report pain ≥ 1 year postoperatively in 30–60% of patients after thoracotomy (141-144) and 20–25% after VATS (141,144). The incidence of taking analgesics is much less (5% after VATS and 20% after thoracotomy) (141,142). Chronic pain has been associated with preoperative narcotic use, the intensity of early postoperative pain and intercostal nerve trauma (145).

The discrepancy between studies investigating QOL and chronic pain is probably due to semantic differences. An earlier review of chronic post-thoracotomy pain found that 50% had some discomfort/pain, ~10% used occasional narcotics, and <5% required more involved treatment (146). Taking this and the more recent studies on QOL and pain together, it appears these rates are still seen after thoracotomy, but approximately half as frequent after VATS.

Nuances and sources of ambiguity Impact of resection margin

Guidelines recommend a resection margin of ≥ 2 cm (from tumor edge to cut lung parenchyma) or a margin to tumor size (M/T) ratio of ≥ 1) (147,148). Clinical practice, however, requires quantification of the risk of a narrow margin so it can be weighed against issues associated with additional resection. The ideal measure is actuarial locoregional recurrence (survival is muddied by unrelated deaths).

Variability in studies of margin distance and M/T ratio (*Tables 8,9*) (53,149-164) likely reflects multiple factors—e.g., adjustment for confounders, proportion of unfit patients or favorable tumors, follow-up duration, resection extent (average margin 15 mm for segment vs. 8 mm for wedge in a prospective study) (165). The data loosely suggest an inflection point around 1 cm, with ~25% recurrence with <1 cm margins. Why Maurizi *et al.* found no difference is unclear (150). The data regarding M/T ratio loosely suggests a locoregional recurrence rate of

1 st author, year (reference)	Years	⊆	Stage	Mean size	Comments	om n/ֈ n	ortion of risk T ª	STA	anent	ə6pə/	x	Outcome	Time period		Margin (mm)	(Li	AVM yc	Factors	lts fidence in
						вэМ		Ά%	S %	W %	N %			220	16-20 11-15	6-10 <5	l giS	îo #	neən
Recurrence															% Recurrence	nce			
Mohiuddin 2014 (149)	01-11	367	cIA1,2	1	Excl BAC	36	+	58	•	100	68	Я	2 yr		9 13	24 29	<.05	6	Σ
Maurizi 2015 (150)	03-13	138	pIA1,2		All hi risk pts	31	+	0	•	100	0	5	۹,	[24] ^b	[25] ^b	[25] ^b	1	ī	
Sienel 2007 (151)	87-02	49	cIA	19		54	+	0	100	ı.	0	5	۹,		م [0]	[23] ^b	ı	ı	
Maurizi 2015 (150)	03-13	182	٩	ı.	All hi risk pts	31	+	0	1	100	0	5	۹,	[25] ^b	[28] ^b	[27] ^b	ı	ī	
Moon 2017 (152)	04-13	39	cIA	17	CTR ≥.5	32	‡	72	26	74	59	Ч	۵ ۱		[18] ^b	[73] ^b	1	ı	
RFS															% 5-year RFS	RES			
Mohiuddin 2014 (149)	01-11	367	cIA1,2		Excl BAC	36	+	58	1	100	68	LR-RFS	2 yr		92 88	80 77	<.05	6	Σ
Dolan 2021 (153)	10-16	695	U	15		51	‡	96	0	100	30	LR-RFS	5 yr		86	82	ı	ı	ı
Maurizi 2015 (150)	03-13	138	plA1,2		All hi risk pts	31	+	0	1	100	0	RFS	5 yr	54	88	53	SN	ω	Σ
Maurizi 2015 (150)	03-13	182	٩		All hi risk pts	31	+	0	ı	100	0	RFS	5 yr	54	48	59	SN	ω	Σ
El-Sherif 2007 (154)	97-04	81	AII-I	21	All hi risk pts	20	+	Some	32	68	•	RFS	5 yr		20	63	•	ı.	
Dolan 2021 (153)	10-16	695	U	15		51	‡	96	ο	100	30	RFS	5 yr		69	65	ı	ı	ı
Wolff ^c 2017 (155)	00-02	138	IA1,2	13	Excl BAC AIS	50	‡	47	T	100	22	RFS	5 yr	ı	87	66	<.05	4	۸L
Moon 2017 (152)	04-13	39	cIA	17	CTR≥.5	32	‡	72	26	74	59	RFS	5 yr		80	24	<.03	13	_
Moon 2017 (152)	04-13	52	cIA1,2 ^d	12	CTR <.5	32	+ + + +	85	35	65	58	RFS	5 yr		100	100	I.	I.	
Masai 2017 (156)	04-13	508	pl-IIA	4	49% AIS MIA	51	+ + + +	0	46	54	1	RFS	5 yr	100	96	74	ı	ı	

מ 5 כ multivariable) 2

events during the study period (in brackets because not an actuarial rate); ^o, 18% of patients from a screening study (I-ELCAD); ^d, 8% cIA3; ^o, staples included in margin measurement; ^t raw incidence of patient population; and policy institutional qualitative estimate from reported proportions of AIS/MIA, low CTR tumors, elective limited resection, invasive tumor size, also used for M/T calculation; ⁹, for entire study (may not be accurate for the subset).

bronchoalveolar carcinoma; f/u, median follow-up (months); hi risk pts, high risk patients (comorbidities precluding lobectomy); L, local recurrence (in same lobe or lobar nodes); LR, locoregional recurrence (in same or adjacent lobe or in intrathoracic nodes); M/T, margin to tumor ratio; MVA, multivariable analysis; Nx, no nodes assessed; BFS, excluded adenocarcinoma in situ; Any R, any recurrence; CTR, consolidation/total tumor ratio of size on CT (lung windows); D Recur, distant recurrence; Excl BAC, ecurrence-free survival; Sig by MVA, statistically significant by multivariable analysis; STAS +/-, spread through air spaces present/absent. AIS,

Table 8 Recurrence outcomes according to margin distance

1 st author, year	Years	c	Stage	Mean	Comments	om n/J	o tion of sk T ^ª	ST	tnemp	ədbə	:	Outcome	Time	Margin/tumor diameter ratio	'tumor er ratio	AVM _V d	actors	aence
				245		nsəM	Prop.	/∧ %	9S %	M %	N %			M/T≥1	M/T <1	ngi2	t to #	fnoð in res
RFS											1			% 5-year RFS	ar RFS			ļ
Sawabata ^e 2012 (157)	99-02	37	HI-I	15	All hi risk pts	>60	+	,	0	100	21	RFS	5 yr	85	53	1		1
Takahashi ^e 2019 (158)		32	HI-I	20	All hi risk pts	39	+	•	28	72	1	RFS	5 yr	92	41	•		
Fernando 2014 (159)	06-10	212	clA	19	All hi risk pts	53	÷	65	31	69	7	L RFS	3 yr	67	66	•	•	•
Tamura 2019 (160)	06-13	141	cl-IIA	23	All hi risk pts	43	‡	53	29	71	~40	RFS	I	Better	Worse	•		•
Moon 2018 (161)	08-15	69	clA1,2	13	Non-lepidic	32	+	88	30	20	Many	RFS	5 yr	97	50	<.04	15	Σ
Moon 2020 (162)	08-17	193	clA1,2	, Ø	Inv size	36	‡ + +	93	48	52	Many	RFS	5 yr	100	77	.03	21	I
Moon 2018 (161)	08-15	64	cIA1,2	11	Lepidic	36	+++++	89	30	70	Many	RFS	5 yr	100	100	•	Т	
Any recurrence														% Recurrence	Irrence			
Schuchert 2007 (163)	02-06	182	HIA	23	All hi risk pts	18	+	37	100	0	Few	Any R	۵ ۱	م [9]	[25] ^b			
Eguchi 2019 (53)	95-14	170	G	10 1	STAS +		+	•	36 9	64 ^g	44 9	Any R	5 yr	59	36	•		1
Eguchi 2019 (53)	95-14	205	cl	10 ¹	STAS –	-	+++	1	36 9	64 ^g	44 9	Any R	5 yr	5	12	•	1	
LR recurrence														% LR recurrence	currence			
El-Sherif 2007 (154)	97-04	81	HIA	21	All hi risk pts	20	+	Some	32	68	1	LR Recur	٦	م [8]	[15] ^b	•		1
Fernando 2014 (159)	06-10	212	cIA	19	All hi risk pts	53	+	65	31	69	7	L Recur	3 yr	14	20	•		1
Eguchi 2019 (53)	95-14	170	G	10 1	STAS +	1	+	1	36 9	64 ^g	44 9	LR Recur	5 yr	16	25	1	1	1
Eguchi 2019 (53)	95-14	205	U	10 ¹	STAS –		+++++++++++++++++++++++++++++++++++++++	1	36 9	64 ^g	44 9	LR Recur	5 yr	0	7	1		
Moon 2018 (161)	08-15	69	cIA1,2	13	Non-lepidic	32	++	88	30	70	Many	LR Recur	۹ ۱	3] ^b	[22] ^b	1	ı.	
Distant recurrence														% D recurrence	urrence			
Eguchi 2019 (53)	95-14	170	5	10 [†]	STAS +	,	+	1	36 °	64 ^e	44 ^e	D Recur	5 yr	13	12	•		
Eguchi 2019 (53)	95-14	205	cl	10 1	STAS –	1	++	1	36 °	64 ^e	44 e	D Recur	5 yr	5	5	•		1
R0 resection														1%	RO			
Sawabata ^e 2004 (164)	99-02	118	cl-IIA	15	All hi risk pts		+	39	100		1	RO	1	100	53	1		

darker = multivariable). For abbreviations, footnotes see legend for *Table* 8.

Journal of Thoracic Disease, Vol 14, No 6 June 2022

Table 9 Recurrence outcomes according to margin to tumor ratio

Detterbeck et al. Evidence for resection extent in healthy patients

~20% for M/T <1 *vs.* ~10% for \geq 1. Margin distance appears to have little impact in primarily GG tumors (152,156).

Most studies have reported whole tumor size. Those reporting invasive size suggest the M/T (invasive) ratio is important (53,162). The discrepancy between the surgeon's and pathologist's margin assessment is another issue (not quantitatively defined). The pathologist typically removes the staple line, and measures the deflated, fixed lung. Studies mostly report the pathologic margin. Surgeons should aim for a surgical margin well beyond a M/T ratio of 1.

In conclusion, for solid tumors evidence loosely suggests a local recurrence rate of ~20–25% for a M/T ratio <1 or a margin <1 cm vs. ~10% for larger margins (recognizing that the pathologic measurement is likely ~3–5 mm less than the surgical assessment).

Impact of STAS

The term "spread through air spaces" (STAS) refers to a microscopic observation of tumor cells adjacent to a lung cancer; the median distance is 1–1.5 mm, but distances of 8–10 mm have been observed (166-169). STAS occurs in essentially all lung cancer types (adenocarcinoma, squamous, small cell, carcinoid, pleomorphic etc.) (169). The reported incidence is quite variable (15–80%) for each tumor type. STAS is rarely observed in adenocarcinoma in situ, minimally invasive adenocarcinoma or pure GG tumors (156,170-174) with some exceptions (29% STAS+ in pure GG, 34% among preinvasive tumors in one study) (175).

STAS is widely associated with worse long-term outcomes (169,176)—but also associated with multiple negative prognostic factors, e.g., aggressive adenocarcinoma subtypes (e.g., solid, micropapillary) (166,167,174,177-181), higher stage (174,175,180,182,183), larger tumors (169,174,175,180-183), and a greater solid component on imaging (172,175,181). No consistent correlation of STAS with genetic characteristics has emerged (169).

In most studies STAS portends worse RFS and higher recurrence rates after sublobar resection (*Tables 10,11*) (156,166-168,170,173,174,178,181-186). This is generally maintained after multivariable adjustment (only limited confounders accounted for). There is less data after lobectomy—STAS portends worse RFS but this is generally not maintained after multivariable adjustment. STAS is associated with a higher distant recurrence rate after sublobar resection in some studies (181,184) but not in others (170,174,186). A greater proportion of favorable tumors doesn't mitigate the negative prognostic impact of STAS.

A simplistic assumption is that STAS represents a

mechanism by which metastasis occurs. This creates a focus on intraoperative detection (frozen-section sensitivity), resection extent and defining a safe margin. However, decades of evidence demonstrate that metastasis is determined by complex cellular transformations, signaling and host-tumor interactions (187-189). STAS may reflect microenvironment evidence of these processes. In other cancers microenvironment evidence of immune recognition of cancer cells and activation of tumor-host interaction predicts long-term outcomes (190). This mental construct suggests that surgical interventions would not affect the impact of STAS.

The available data is inconclusive whether a negative prognostic impact of STAS can be altered by a more extensive resection. Few studies have addressed this with conflicting results (Table 11) (53,178,185). In an extensively adjusted retrospective analysis Eguchi et al. found that if STAS is present, lobectomy is associated with better RFS and fewer recurrences than sublobar resection (53). Eguchi et al. also observed that recurrences after sublobar resection in STAS + tumors were associated with an M/T ratio of <1 (this margin/STAS analysis was unadjusted for any confounders) (53). The observation invited speculation that a wider margin might mitigate the negative prognostic impact of STAS. Another unadjusted analysis of sublobar resection found that STAS was associated with a similar increase in loco-regional recurrence for $M/T \ge 1$ as for M/T <1 (174).

Single vs. multi-segmentectomy

A right upper lobectomy is arguably the same as a left upper tri-segmentectomy, and a right middle lobectomy the same as lingulectomy. In database studies the proportion of such "lobe-like" segmentectomies is unavailable. In single-institution series, the proportion is 20–40% (43,46,51,191,192), and 30–55% of segmentectomies involve \geq 3 segments (43,46,51,191,192). Studies involving many multi-segmentectomies found no OS or LCSS difference between segmentectomy vs. lobectomy (43,46,51).

Anatomic location

Whether the tumor size and anatomic location confidently permit an adequate margin is important in deciding the resection extent in an individual patient. Wedge resection is only feasible for tumors in the outer third of the lung (from the pleural space to the hilum). Achieving an adequate margin is difficult even for segmentectomy when tumors are central or near an intersegmental boundary. A simulation model estimated that ~25–33% of 1–2 cm tumors would

1 st outbook wood				acold	tu	⊥ _P ou oţ	B		MVA			Time	Suble	Sublobar resection	tion		Lobectomy	
r autrol, year (reference)	Years	ة Z	Stage	size ^a	მოობე	Proporti Iow risk	SAT2 %	_в хN %	# of factors	əbitnoƏ IuzəЯ ni	Outcome	period	STAS –	STAS +	Sig by MVA	STAS -	STAS +	Sig by MVA
RFS															% 5-year RFS	ar RFS		
Yanagawa 2018 (168)	00-14	80/40	pl-IIA		Squam	ċ	20/20	1	1	1	RFS	5 yr	61	19	1	71	48	
Kadota 2017 (184)	99-12	92/42	AII-I	ı.	Squam	ċ	35/33	•	1	i.	RFS	5 yr	99	39	I	70	63	•
Kagimoto 2021 (185)	07-20	348/261	cIA °	20/14	Ad, Seg	+	48	Few	9	_	RFS	5 yr	93	81	1	06	68	ı.
Ren 2019 (166)	10-12	634/118	plA	ı.	Pd	‡	29/36	•	7	٨L	RFS	5 yr	92	67	<.001	88	81	PS d
Shiono 2018 (182)	04-17	329/185	cIA	19/16	1	‡	22/17	•	13	٨L	RFS	5 yr	82	54	<.02	91	20	NS
Han 2021 (174)	11-18	648/222	cIA	ı.	PA	‡	32/15	1	6	Σ	RFS	5 yr	66	63	.001	97	62	.02
Toyokawa 2018 (183)	03-12	185/89	pl-II	T	PA	+	64/38	1	13	٨L	RFS	5 yr	97	66 ^f	•	94	22	1
Uruga 2017 (173)	03-09	163/45	pIA1,2	T	Ad	~	54/24	1	₽	_	RFS	5 yr	96 [°]	83 [°]	NS	100 ^e	87 ^e	SN
Toyokawa 2018 (186)	03-12	-/82	ll-lq	ı	Ad	‡	-/38	1	÷	٨L	RFS	5 yr	97	69	<.01	ı	ı	ŀ
Chae 2021 (181)	09-16	-/115	cIA	ı	Ad	‡ ‡	-/17	,	ı	I	RFS	5 yr	98	59	.001	86	84	T
Masai 2017 (156)	04-13	-/508	pl-IIA	14		+ + + +	-/15		1	1	RFS	5 yr	97	86	-	I	I	1
Any recurrence															% Recurrence	rrence		
Kadota 2015 (167)	92-06	291/120	pIA1,2	[15] 9	PA	+	37 /38	0/43	œ	٨L	Any R	5 yr	ŧ	43	<.02	10	13	
Shiono 2020 (170)	04-18	-/100	cIA	10 9	Wedge	+	17	93 ^g	15	Σ	Any R	5 yr	34	57	<.03	ı	1	
Shiono 2020 (170)	04-18	-/117	cIA	е _н	Seg	‡	15	0/0	15	Σ	Any R	5 yr	ø	33	NS	I	1	ı.
Shiono 2020 (170)	04-18	-/117	cIA	4 L	1	+	15	0/93	1		Any R	-	[13]	[35] ⁻	I	I	I	I
Han 2021 (174)	11-18	648/222	cIA	ı	Ad	‡	32/15	1	ı	I	Any R		-	10	ı	2	10	ı
Kadota 2019 (178)	99-13	376/114	ō	ı	Ad	‡	1		ī	T	Any R	5 yr	2	52	1	2	34	1
Toyokawa 2018 (186)	03-12	-/82	ll-Id	1	PA	‡	-/38		1	T	Any R		6	[29]	1	I	ı	1
Chae 2021 (181)	09-16	-/115	cIA	1	PA	+ + +	-/17	1	T	ı.	Any R		ო	40	-001	I	I	ı.
Masai 2017 (156)	04-13	-/508	pl-IIA	14	1	+ + + +	-/15	•	7	_	Any R	5 yr	Ш	П	NS	'	I	'
Loco-regional recurrence	ence													% L	oco-regior	% Loco-regional recurrence	ce	
Kadota 2015 (167)	92-06	-/120	pIA1,2	[15] ^g	PA	+	-/38	-/43	ı	Т	LR Recur	5 yr	4	22	T	I	1	т
Kadota 2019 (178)	99-13	-/114	U	ı	Ad	‡	T	1	ı	T	LR Recur	5 yr	-	43	ı	ı	ı	ŀ
Shiono 2020 (170)	04-18	-/117	cIA	۲ ^հ	ı	‡	15	0/93	ı	Ţ	LR Recur	-,	6	[26] ⁻	1	ı	ı	T
Han 2021 (174)	11-18	648/222	cIA	ı	Рd	‡	32/15		ı	T	LR Recur	-	0	7	1	-	4	T
Toyokawa 2018 (186)	03-12	-/82	ll-Id	T	Ad	‡	-/38	,	ı	I	LR Recur		5	[26]	ı	I	I	T
Chae 2021 (181)	09-16	-/115	cIA	1	PA	‡ ‡ +	-/17	1	1		LR Recur		-	25	•	I	I	1
Masai 2017 (156)	04-13	-/508	pl-IIA	14		‡	-/15	•	7	_	LR Recur	5 yr	1	HR 3.14	<.04	ı	1	·

otrat of a company and commence biobot note of the		I ~ - J ~ ~ mm																
1 st author, year (reference)	Years	e Z	Stage	Mean size	stnem	ortion of isk T ^b	ⁱⁱ ədc	را		adence of A #	stius	Outcome	Time	S	STAS –		STAS +	
					ლიე		רי %	IS %	ы % Б	factors Conf				SL I	Lobe ^{Sig by} MVA	A SL	Lobe	Sig by MVA
LCSS															% 5.	% 5-year LCSS		
Eguchi 2019 (53)	95-14	422/276	ט	11 7	Ad	‡	50	50	44	19		LCSS	5 yr	96	96 NS	84	92	.02
RFS															3 %	% 5-year RFS		
Kagimoto 2021 (185)	07-20	348/261	cIA ^k	15/15	Ad Seg	+	63	37 F	Few	9		RFS	5 yr	1	 	83	75	NS
Any recurrence															% An	% Any recurrence	e	
Kagimoto 2021 (185)	07-20	348/261	cIA ^k	15/15	Ad Seg	+	63	37 F	Few	9		Any R	1	ı	1	4	13	<.04
Kadota 2019 (178)	99-13	353/137	0	ı	Ad	‡	77	23	1	1		Any R	5 yr	0	2	52	8	ı
Eguchi 2019 (53)	95-14	422/276	cl	11 "	Ad	‡	50	50	44	19 1	н	Any R	5 yr	6	6 NS	39	16	<.001
Loco-regional recurrence	rence														% Loco-re	% Loco-regional recurrence	Irrence	
Kagimoto 2021 (185)	07-20	348/261	cIA ^k	15/15	Ad Seg	+	63	37 F	Few	1	-	LR Recur	ı	1	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	2	œ	1
Kadota 2019 (178)	99-13	-/137	cl	1	Ad	‡	77	23	-	-	-	LR Recur	5 yr	I	1	43	23	I
Distant recurrence															% Dist	% Distant recurrence	nce	
Kagimoto 2021 (185)	07-20	348/261	cIA ^k	15/15	Ad Seg	+	63	37 F	Few			D Recur	ı	1	1	ო	13	T
Kadota 2019 (178)	99-13	-/137	cl	1	Ad	‡	77	23	-			D Recur	5 yr	I	1	32	19	I
Inclusion criteria (Tables 10,11): studies 2000-2021 reporting on STAS relative to resection extent (sublobar vs. lobectomy), >50 patients. Bold highlights better outcome	ables 10	,11): studi	ies 200(0-2021 r	eporting	on STA	S relativ	ve to re	esectio	n extent	(sublo	bar vs. lot	pectomy),	≥50 patie	ints. Bold h	nighlights	better ou	utcome
(>2-point difference); Light green shading highlights statistically significant difference favoring lobectomy (lighter shade = univariable; darker = multivariable); pink highlights	i); Light (green shau	ding hig	hlights s	statistically	/ signific	cant dif	ference	Favori	ing lobeci	tomy (I	ighter sha	de = univ	ariable; dá	arker = mul	tivariable); pink hig	Ihlights
statistically significant adjusted difference favoring sublobar resection.	int adjus:	ted differe.	ince favo	oring sub	olobar rese	∋ction.												
^a . reported by cohorts: lobe/sublobar: ^b . gualitative	rts: lobe/	/sublobar;	b. auali	tative es	timate fro	m repor	ted pro	portior	Is of A	IS/MIA. Ic	W CTF	3 tumors.	elective li	nited rese	estimate from reported proportions of AIS/MIA. low CTB tumors. elective limited resection. institutional policy and patient	utional p	olicy and	patient

, comparing high STAS to no STAS cohorts; ^f, many of the STAS+ patients were used for M/T calculation;¹, raw incidence of events during the study period (in brackets because not an actuarial rate);¹, total for entire study cohort; ^k, assessed by invasive compromised patients who underwent wedge resections and suffered unrelated deaths; ⁹, for entire study (may not be accurate for the subset); ⁿ, invasive tumor size, also population, clinical trial participation (JCOG 0802); $^{\circ}$, invasive tumor size; d , P=0.057; ŝ tumor size.

Ad, adenocarcinoma; Any R, any recurrence; CTR, consolidation/total tumor ratio of size on CT (lung windows); D Recur, distant recurrence; HR, hazard ratio; LCSS, lung cancer specific survival; Lobe, lobectomy; LR Recur, locoregional recurrence (in same or adjacent lobe or in intrathoracic nodes); MVA, multivariable analysis; NS, not segmentectomy; SL, sublobar resection; Squam, squamous carcinoma; Sig by MVA, statistically significant by multivariable analysis; STAS +/-, spread through air spaces present/absent; T, tumor; yr, year. no nodes assessed; RFS, recurrence-free survival; Seg, significant (P>0.05); Nx,

be amenable to segmentectomy (defined as ≥ 2 cm from an intersegmental plane); for bi-segmentectomy ~50% would meet this criterion (assuming uniform tumor distribution throughout the lungs) (193).

Summary of outcomes in healthy patients

In healthy patients contemporary RCTs demonstrate equivalent perioperative mortality for segmentectomy or wedge vs. lobectomy (1–4% 90-day mortality). The incidence of major complications is also low (5–15% grade \geq 3) and not improved by sublobar resection. A significant benefit to VATS over thoracotomy has been demonstrated extensively for lobectomy; this also appears true for segmentectomy. Pain and impaired QOL is generally resolved by 3 months after VATS resection.

Adjusted NRCs with high confidence that results reflect the treatment demonstrate worse OS for segmentectomy or wedge resection than lobectomy. Multiple additional NRCs with greater residual confounding mostly favor lobectomy; statistical significance is fairly consistent for OS and LCSS for wedge but less so for segmentectomy *vs.* lobectomy. While we await mature results from RCTs, the aggregate evidence indicates meaningfully worse long-term outcomes after segmentectomy or wedge resection than lobectomy in healthy patients with cI NSCLC.

VATS resection has little long-term impact on QOL, but open resection results in persistently worse QOL. A QOL benefit to sublobar resection is unclear due to confounding by VATS/open approach. Sublobar resection may attenuate an increase in dyspnea that is commonly noted after lobectomy. However, PFTs demonstrate no meaningful advantage for segmentectomy over lobectomy in healthy patients, particularly when including multisegmentectomies.

Evidence suggests no meaningful difference in short-, intermediate- or long-term outcomes for a "lobe-like" multi-segmentectomy vs. lobectomy. The risk of an inadequate margin given an individual tumor's anatomic location is an important consideration. Locoregional recurrence rates of ~20–25% for margins of <1 cm or a margin/tumor ratio of <1 are half as frequent with larger margins for solid tumors; margin appears to have less impact in primarily GG tumors. Worse long-term outcomes are reported when STAS is present (especially after sublobar resection); this is confounded because STAS is associated with many negative prognostic factors. It is unclear whether the impact of STAS can be mitigated by converting to a lobectomy.

Short-term and long-term outcomes for segmentectomy or wedge resection *vs.* lobectomy are summarized in Table S2-2. A benefit or detriment is qualitatively depicted relative to clinically meaningful differences, together with the confidence in and consistency of the evidence. This provides a succinct summary that can inform judgment for individual patients, as discussed in the Part 1 paper (1).

Conclusions

Choosing which type of resection is best for a particular patient demands balancing various factors and outcomes. This analysis of the relevant evidence in generally healthy patients provides a foundation for a framework to facilitate individualized decision-making across the spectrum of lung cancer patients.

Acknowledgments

Funding: None.

Footnote

Provenance and Peer Review: This article was commissioned by the editorial office, *Journal of Thoracic Disease* for the series "A Guide for Managing Patients with Stage I NSCLC: Deciding between Lobectomy, Segmentectomy, Wedge, SBRT and Ablation". The article has undergone external peer review.

Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at https://jtd.amegroups.com/article/view/10.21037/jtd-21-1824/coif). The series "A Guide for Managing Patients with Stage I NSCLC: Deciding between Lobectomy, Segmentectomy, Wedge, SBRT and Ablation" was commissioned by the editorial office without any funding or sponsorship. FCD served as the unpaid Guest Editor of the series. HSP serves as an unpaid editorial board member of *Journal of Thoracic Disease*. HSP reports research funding from RefleXion Medical; consulting fees from AstraZeneca; honoraria and speaking fees from Galera Therapeutics; all unrelated to current work. The authors have no other conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related

Detterbeck et al. Evidence for resection extent in healthy patients

to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

Open Access Statement: This is an Open Access article distributed in accordance with the Creative Commons Attribution-NonCommercial-NoDerivs 4.0 International License (CC BY-NC-ND 4.0), which permits the non-commercial replication and distribution of the article with the strict proviso that no changes or edits are made and the original work is properly cited (including links to both the formal publication through the relevant DOI and the license). See: https://creativecommons.org/licenses/by-nc-nd/4.0/.

References

- Detterbeck FC, Blasberg JD, Woodard GA, et al. A guide for managing patients with stage I NSCLC: deciding between lobectomy, segmentectomy, wedge, SBRT and ablation—part 1: a guide to decision-making. J Thorac Dis 2022. doi: 10.21037/jtd-21-1823
- Bade BC, Blasberg JD, Mase VJ Jr, et al. A guide for managing patients with stage I NSCLC: deciding between lobectomy, segmentectomy, wedge, SBRT and ablation part 3: systematic review of evidence regarding surgery in compromised patients or specific tumors. J Thorac Dis 2022. doi: 10.21037/jtd-21-1825
- Park HS, Detterbeck FC, Madoff DC, et al. A guide for managing patients with stage I NSCLC: deciding between lobectomy, segmentectomy, wedge, SBRT and ablation part 4: systematic review of evidence involving SBRT and ablation. J Thorac Dis 2022. doi: 10.21037/jtd-21-1826
- Elze MC, Gregson J, Baber U, et al. Comparison of Propensity Score Methods and Covariate Adjustment: Evaluation in 4 Cardiovascular Studies. J Am Coll Cardiol 2017;69:345-57.
- Hade EM, Lu B. Bias associated with using the estimated propensity score as a regression covariate. Stat Med 2014;33:74-87.
- Benedetto U, Head SJ, Angelini GD, et al. Statistical primer: propensity score matching and its alternatives. Eur J Cardiothorac Surg 2018;53:1112-7.
- Sterne JA, Hernán MA, Reeves BC, et al. ROBINS-I: a tool for assessing risk of bias in non-randomised studies of interventions. BMJ 2016;355:i4919.
- Ginsberg RJ, Rubinstein LV. Randomized trial of lobectomy versus limited resection for T1 N0 non-small cell lung cancer. Lung Cancer Study Group. Ann Thorac Surg 1995;60:615-22; discussion 622-3.

- Altorki NK, Wang X, Wigle D, et al. Perioperative mortality and morbidity after sublobar versus lobar resection for early-stage non-small-cell lung cancer: post-hoc analysis of an international, randomised, phase 3 trial (CALGB/Alliance 140503). Lancet Respir Med 2018;6:915-24.
- Suzuki K, Saji H, Aokage K, et al. Comparison of pulmonary segmentectomy and lobectomy: Safety results of a randomized trial. J Thorac Cardiovasc Surg 2019;158:895-907.
- 11. Stamatis G, Leschber G, Schwarz B, et al. Perioperative course and quality of life in a prospective randomized multicenter phase III trial, comparing standard lobectomy versus anatomical segmentectomy in patients with nonsmall cell lung cancer up to 2 cm, stage IA (7th edition of TNM staging system). Lung Cancer 2019;138:19-26.
- Linden PA, D'Amico TA, Perry Y, et al. Quantifying the safety benefits of wedge resection: a society of thoracic surgery database propensity-matched analysis. Ann Thorac Surg 2014;98:1705-11; discussion 1711-2.
- Stokes WA, Bronsert MR, Meguid RA, et al. Post-Treatment Mortality After Surgery and Stereotactic Body Radiotherapy for Early-Stage Non-Small-Cell Lung Cancer. J Clin Oncol 2018;36:642-51.
- Zhang Z, Feng H, Zhao H, et al. Sublobar resection is associated with better perioperative outcomes in elderly patients with clinical stage I non-small cell lung cancer: a multicenter retrospective cohort study. J Thorac Dis 2019;11:1838-48.
- Bédat B, Abdelnour-Berchtold E, Perneger T, et al. Comparison of postoperative complications between segmentectomy and lobectomy by video-assisted thoracic surgery: a multicenter study. J Cardiothorac Surg 2019;14:189.
- Khullar OV, Liu Y, Gillespie T, et al. Survival After Sublobar Resection versus Lobectomy for Clinical Stage IA Lung Cancer: An Analysis from the National Cancer Data Base. J Thorac Oncol 2015;10:1625-33.
- Tsutani Y, Tsubokawa N, Ito M, et al. Postoperative complications and prognosis after lobar resection versus sublobar resection in elderly patients with clinical Stage I non-small-cell lung cancer. Eur J Cardiothorac Surg 2018;53:366-71.
- Husain ZA, Kim AW, Yu JB, et al. Defining the High-Risk Population for Mortality After Resection of Early Stage NSCLC. Clin Lung Cancer 2015;16:e183-7.
- 19. Shirvani SM, Jiang J, Chang JY, et al. Lobectomy, sublobar resection, and stereotactic ablative radiotherapy for early-

2378

Journal of Thoracic Disease, Vol 14, No 6 June 2022

stage non-small cell lung cancers in the elderly. JAMA Surg 2014;149:1244-53.

- Okami J, Shintani Y, Okumura M, et al. Demographics, Safety and Quality, and Prognostic Information in Both the Seventh and Eighth Editions of the TNM Classification in 18,973 Surgical Cases of the Japanese Joint Committee of Lung Cancer Registry Database in 2010. J Thorac Oncol 2019;14:212-22.
- 21. Thai AA, Stuart E, Te Marvelde L, et al. Hospital lung surgery volume and patient outcomes. Lung Cancer 2019;129:22-7.
- 22. Fagundes CP, Shi Q, Vaporciyan AA, et al. Symptom recovery after thoracic surgery: Measuring patientreported outcomes with the MD Anderson Symptom Inventory. J Thorac Cardiovasc Surg 2015;150:613-9.e2.
- Detterbeck F, Antonicelli A, Okada M. Results of Video-Assisted Techniques for Resection of Lung Cancer. In: Pass H, Ball D, Scagliotti G. editors. Thoracic Oncology: The IASLC Multidisciplinary Approach (2nd Edition): IASLC, 2018.
- 24. Bendixen M, Jørgensen OD, Kronborg C, et al. Postoperative pain and quality of life after lobectomy via video-assisted thoracoscopic surgery or anterolateral thoracotomy for early stage lung cancer: a randomised controlled trial. Lancet Oncol 2016;17:836-44.
- 25. Smith CB, Kale M, Mhango G, et al. Comparative outcomes of elderly stage I lung cancer patients treated with segmentectomy via video-assisted thoracoscopic surgery versus open resection. J Thorac Oncol 2014;9:383-9.
- 26. Ghaly G, Kamel M, Nasar A, et al. Video-Assisted Thoracoscopic Surgery Is a Safe and Effective Alternative to Thoracotomy for Anatomical Segmentectomy in Patients With Clinical Stage I Non-Small Cell Lung Cancer. Ann Thorac Surg 2016;101:465-72; discussion 472.
- Schuchert MJ, Pettiford BL, Pennathur A, et al. Anatomic segmentectomy for stage I non-small-cell lung cancer: comparison of video-assisted thoracic surgery versus open approach. J Thorac Cardiovasc Surg 2009;138:1318-25.e1.
- Nomori H, Cong Y, Sugimura H. Limited thoracotomy for segmentectomy: a comparison of postoperative pain with thoracoscopic lobectomy. Surg Today 2016;46:1243-8.
- Handa Y, Tsutani Y, Mimae T, et al. Surgical Outcomes of Complex Versus Simple Segmentectomy for Stage I Non-Small Cell Lung Cancer. Ann Thorac Surg 2019;107:1032-9.
- 30. Detterbeck FC. Lobectomy versus limited resection in T1N0 lung cancer. Ann Thorac Surg 2013;96:742-4.
- 31. Lederle FA. Lobectomy versus limited resection in T1 N0

lung cancer. Ann Thorac Surg 1996;62:1249-50.

- 32. Yang F, Sui X, Chen X, et al. Sublobar resection versus lobectomy in Surgical Treatment of Elderly Patients with early-stage non-small cell lung cancer (STEPS): study protocol for a randomized controlled trial. Trials 2016;17:191.
- 33. Tanaka K, Tsutani Y, Wakabayashi M, et al. Sublobar resection versus lobectomy for patients with resectable stage I non-small cell lung cancer with idiopathic pulmonary fibrosis: a phase III study evaluating survival (JCOG1708, SURPRISE). Jpn J Clin Oncol 2020;50:1076-9.
- 34. Asamura H, Okada M, Saji H, et al. Randomized Trial of Segmentectomy Compared to Lobectomy in Small-Sized Peripheral Non-Small Cell Lung Cancer. AATS Virtual 2021.
- 35. Saji H, Okada M, Tsuboi M, et al. Segmentectomy versus lobectomy in small-sized peripheral non-small-cell lung cancer (JCOG0802/WJOG4607L): a multicentre, openlabel, phase 3, randomised, controlled, non-inferiority trial. Lancet 2022;399:1607-17.
- 36. Cao J, Yuan P, Wang Y, et al. Survival Rates After Lobectomy, Segmentectomy, and Wedge Resection for Non-Small Cell Lung Cancer. Ann Thorac Surg 2018;105:1483-91.
- Onaitis MW, Furnary AP, Kosinski AS, et al. Equivalent Survival Between Lobectomy and Segmentectomy for Clinical Stage IA Lung Cancer. Ann Thorac Surg 2020;110:1882-91.
- 38. Li F, Zhao Y, Yuan L, et al. Oncologic outcomes of segmentectomy vs lobectomy in pathologic stage IA (≤2 cm) invasive lung adenocarcinoma: A population-based study. J Surg Oncol 2020;121:1132-9.
- Koike T, Kitahara A, Sato S, et al. Lobectomy Versus Segmentectomy in Radiologically Pure Solid Small-Sized Non-Small Cell Lung Cancer. Ann Thorac Surg 2016;101:1354-60.
- 40. Zhao ZR, Situ DR, Lau RWH, et al. Comparison of Segmentectomy and Lobectomy in Stage IA Adenocarcinomas. J Thorac Oncol 2017;12:890-6.
- Moon MH, Moon YK, Moon SW. Segmentectomy versus lobectomy in early non-small cell lung cancer of 2 cm or less in size: A population-based study. Respirology 2018;23:695-703.
- Yendamuri S, Sharma R, Demmy M, et al. Temporal trends in outcomes following sublobar and lobar resections for small (≤ 2 cm) non-small cell lung cancers--a Surveillance Epidemiology End Results database analysis. J Surg Res

Detterbeck et al. Evidence for resection extent in healthy patients

2380

2013;183:27-32.

- Yamashita S, Tokuishi K, Anami K, et al. Thoracoscopic segmentectomy for T1 classification of non-small cell lung cancer: a single center experience. Eur J Cardiothorac Surg 2012;42:83-8.
- 44. Qu X, Wang K, Zhang T, et al. Long-term outcomes of stage I NSCLC (≤3 cm) patients following segmentectomy are equivalent to lobectomy under analogous extent of lymph node removal: a PSM based analysis. J Thorac Dis 2017;9:4561-73.
- 45. Chan EG, Chan PG, Mazur SN, et al. Outcomes with segmentectomy versus lobectomy in patients with clinical T1cN0M0 non-small cell lung cancer. J Thorac Cardiovasc Surg 2021;161:1639-1648.e2.
- 46. Landreneau RJ, Normolle DP, Christie NA, et al. Recurrence and survival outcomes after anatomic segmentectomy versus lobectomy for clinical stage I nonsmall-cell lung cancer: a propensity-matched analysis. J Clin Oncol 2014;32:2449-55.
- 47. Fan X, Liang Y, Bai Y, et al. Conditional survival rate estimates of lobectomy, segmentectomy and wedge resection for stage IA1 non-small cell lung cancer: A population-based study. Oncol Lett 2020;20:1607-18.
- 48. Dai C, Shen J, Ren Y, et al. Choice of Surgical Procedure for Patients With Non-Small-Cell Lung Cancer ≤ 1 cm or > 1 to 2 cm Among Lobectomy, Segmentectomy, and Wedge Resection: A Population-Based Study. J Clin Oncol 2016;34:3175-82.
- Whitson BA, Groth SS, Andrade RS, et al. Survival after lobectomy versus segmentectomy for stage I non-small cell lung cancer: a population-based analysis. Ann Thorac Surg 2011;92:1943-50.
- Dziedzic R, Zurek W, Marjanski T, et al. Stage I nonsmall-cell lung cancer: long-term results of lobectomy versus sublobar resection from the Polish National Lung Cancer Registry. Eur J Cardiothorac Surg 2017;52:363-9.
- 51. Hwang Y, Kang CH, Kim HS, et al. Comparison of thoracoscopic segmentectomy and thoracoscopic lobectomy on the patients with non-small cell lung cancer: a propensity score matching study. Eur J Cardiothorac Surg 2015;48:273-8.
- 52. Razi SS, Nguyen D, Villamizar N. Lobectomy does not confer survival advantage over segmentectomy for nonsmall cell lung cancer with unsuspected nodal disease. J Thorac Cardiovasc Surg 2020;159:2469-2483.e4.
- 53. Eguchi T, Kameda K, Lu S, et al. Lobectomy Is Associated with Better Outcomes than Sublobar Resection in Spread through Air Spaces (STAS)-Positive T1 Lung

Adenocarcinoma: A Propensity Score-Matched Analysis. J Thorac Oncol 2019;14:87-98.

- 54. Yu Y, Huang R, Wang P, et al. Sublobectomy versus lobectomy for long-term survival outcomes of earlystage non-small cell lung cancer with a tumor size ≤2 cm accompanied by visceral pleural invasion: a SEER population-based study. J Thorac Dis 2020;12:592-604.
- 55. Eguchi T, Bains S, Lee MC, et al. Impact of Increasing Age on Cause-Specific Mortality and Morbidity in Patients With Stage I Non-Small-Cell Lung Cancer: A Competing Risks Analysis. J Clin Oncol 2017;35:281-90.
- 56. Liang Y, Fan X, Bai Y, et al. Conditional survival analysis of four treatment strategies for patients with stage I non-small cell lung cancer. Oncol Lett 2019;18:1089-98.
- 57. Dolan DP, White A, Mazzola E, et al. Outcomes of superior segmentectomy versus lower lobectomy for superior segment Stage I non-small-cell lung cancer are equivalent: An analysis of 196 patients at a single, high volume institution. J Surg Oncol 2021;123:570-8.
- Boyer MJ, Williams CD, Harpole DH, et al. Improved Survival of Stage I Non-Small Cell Lung Cancer: A VA Central Cancer Registry Analysis. J Thorac Oncol 2017;12:1814-23.
- Speicher PJ, Gu L, Gulack BC, et al. Sublobar Resection for Clinical Stage IA Non-small-cell Lung Cancer in the United States. Clin Lung Cancer 2016;17:47-55.
- Subramanian M, McMurry T, Meyers BF, et al. Long-Term Results for Clinical Stage IA Lung Cancer: Comparing Lobectomy and Sublobar Resection. Ann Thorac Surg 2018;106:375-81.
- 61. Cox ML, Yang CJ, Speicher PJ, et al. The Role of Extent of Surgical Resection and Lymph Node Assessment for Clinical Stage I Pulmonary Lepidic Adenocarcinoma: An Analysis of 1991 Patients. J Thorac Oncol 2017;12:689-96.
- 62. Nakamura H, Taniguchi Y, Miwa K, et al. Comparison of the surgical outcomes of thoracoscopic lobectomy, segmentectomy, and wedge resection for clinical stage I non-small cell lung cancer. Thorac Cardiovasc Surg 2011;59:137-41.
- 63. Smith CB, Swanson SJ, Mhango G, et al. Survival after segmentectomy and wedge resection in stage I non-smallcell lung cancer. J Thorac Oncol 2013;8:73-8.
- 64. Koike T, Koike T, Yoshiya K, et al. Risk factor analysis of locoregional recurrence after sublobar resection in patients with clinical stage IA non-small cell lung cancer. J Thorac Cardiovasc Surg 2013;146:372-8.
- 65. Zhang Y, Sun Y, Chen H. A propensity score matching analysis of survival following segmentectomy or wedge

Journal of Thoracic Disease, Vol 14, No 6 June 2022

resection in early-stage lung invasive adenocarcinoma or squamous cell carcinoma. Oncotarget 2016;7:13880-5.

- 66. Zhao M, Lu T, Huang Y, et al. Survival and Long-Term Cause-Specific Mortality Associated With Stage IA Lung Adenocarcinoma After Wedge Resection vs. Segmentectomy: A Population-Based Propensity Score Matching and Competing Risk Analysis. Front Oncol 2019;9:5593.
- 67. Stiles BM, Mao J, Harrison S, et al. Extent of lymphadenectomy is associated with oncological efficacy of sublobar resection for lung cancer ≤2 cm. J Thorac Cardiovasc Surg 2019;157:2454-2465.e1.
- 68. Zhang B, Liu R, Ren D, et al. Comparison of Lobectomy and Sublobar Resection for Stage IA Elderly NSCLC Patients (≥70 Years): A Population-Based Propensity Score Matching's Study. Front Oncol 2021;11:610638.
- Cao C, Gupta S, Chandrakumar D, et al. Meta-analysis of intentional sublobar resections versus lobectomy for early stage non-small cell lung cancer. Ann Cardiothorac Surg 2014;3:134-41.
- 70. Cao C, Chandrakumar D, Gupta S, et al. Could less be more?-A systematic review and meta-analysis of sublobar resections versus lobectomy for non-small cell lung cancer according to patient selection. Lung Cancer 2015;89:121-32.
- Bao F, Ye P, Yang Y, et al. Segmentectomy or lobectomy for early stage lung cancer: a meta-analysis. Eur J Cardiothorac Surg 2014;46:1-7.
- Bedetti B, Bertolaccini L, Rocco R, et al. Segmentectomy versus lobectomy for stage I non-small cell lung cancer: a systematic review and meta-analysis. J Thorac Dis 2017;9:1615-23.
- 73. Guo J, Liu Y, Tian X, et al. Less is more in solid-dominant lung cancer? Sublobar resection versus lobectomy for solid-dominant stage IA non-small-cell lung cancer: A meta-analysis study. Mol Clin Oncol 2019;11:465-73.
- Hennon M, Landreneau RJ. Role of Segmentectomy in Treatment of Early-Stage Non-Small Cell Lung Cancer. Ann Surg Oncol 2018;25:59-63.
- 75. Rao S, Ye L, Min L, et al. Meta-analysis of segmentectomy versus lobectomy for radiologically pure solid or solid-dominant stage IA non-small cell lung cancer. J Cardiothorac Surg 2019;14:197.
- Huang CS, Hsu PK, Chen CK, et al. Surgeons' preference sublobar resection for stage I NSCLC less than 3 cm. Thorac Cancer 2020;11:907-17.
- 77. Kamigaichi A, Tsutani Y, Mimae T, et al. Prognosis of segmentectomy and lobectomy for radiologically

aggressive small-sized lung cancer. Eur J Cardiothorac Surg 2020;58:1245-53.

- El-Sherif A, Gooding WE, Santos R, et al. Outcomes of sublobar resection versus lobectomy for stage I non-small cell lung cancer: a 13-year analysis. Ann Thorac Surg 2006;82:408-15; discussion 415-6.
- Tsutani Y, Handa Y, Shimada Y, et al. Comparison of cancer control between segmentectomy and wedge resection in patients with clinical stage IA non-small cell lung cancer. J Thorac Cardiovasc Surg 2021;162:1244-1252.e1.
- Altorki NK, Kamel MK, Narula N, et al. Anatomical Segmentectomy and Wedge Resections Are Associated with Comparable Outcomes for Patients with Small cT1N0 Non-Small Cell Lung Cancer. J Thorac Oncol 2016;11:1984-92.
- Yoshikawa K, Tsubota N, Kodama K, et al. Prospective study of extended segmentectomy for small lung tumors: the final report. Ann Thorac Surg 2002;73:1055-8; discussion 1058-9.
- Takizawa T, Haga M, Yagi N, et al. Pulmonary function after segmentectomy for small peripheral carcinoma of the lung. J Thorac Cardiovasc Surg 1999;118:536-41.
- Harada H, Okada M, Sakamoto T, et al. Functional advantage after radical segmentectomy versus lobectomy for lung cancer. Ann Thorac Surg 2005;80:2041-5.
- Kashiwabara K, Sasaki J, Mori T, et al. Relationship between functional preservation after segmentectomy and volume-reduction effects after lobectomy in stage I nonsmall cell lung cancer patients with emphysema. J Thorac Oncol 2009;4:1111-6.
- 85. Yoshimoto K, Nomori H, Mori T, et al. Quantification of the impact of segmentectomy on pulmonary function by perfusion single-photon-emission computed tomography and multidetector computed tomography. J Thorac Cardiovasc Surg 2009;137:1200-5.
- Saito H, Nakagawa T, Ito M, et al. Pulmonary function after lobectomy versus segmentectomy in patients with stage I non-small cell lung cancer. World J Surg 2014;38:2025-31.
- Nomori H, Cong Y, Sugimura H. Systemic and regional pulmonary function after segmentectomy. J Thorac Cardiovasc Surg 2016;152:747-53.
- Suzuki H, Morimoto J, Mizobuchi T, et al. Does segmentectomy really preserve the pulmonary function better than lobectomy for patients with early-stage lung cancer? Surg Today 2017;47:463-9.
- 89. Gu Z, Wang H, Mao T, et al. Pulmonary function changes

Detterbeck et al. Evidence for resection extent in healthy patients

after different extent of pulmonary resection under videoassisted thoracic surgery. J Thorac Dis 2018;10:2331-7.

- 90. Tane S, Nishio W, Fujibayashi Y, et al. Thoracoscopic left S1+2 segmentectomy as a good resolution for preserving pulmonary function. Interact Cardiovasc Thorac Surg 2020;31:331-8.
- 91. Keenan RJ, Landreneau RJ, Maley RH Jr, et al. Segmental resection spares pulmonary function in patients with stage I lung cancer. Ann Thorac Surg 2004;78:228-33; discussion 228-33.
- Nomori H, Mori T, Ikeda K, et al. Segmentectomy for selected cT1N0M0 non-small cell lung cancer: a prospective study at a single institute. J Thorac Cardiovasc Surg 2012;144:87-93.
- 93. Nomori H, Shiraishi A, Cong Y, et al. Differences in postoperative changes in pulmonary functions following segmentectomy compared with lobectomy. Eur J Cardiothorac Surg 2018;53:640-7.
- 94. Macke RA, Schuchert MJ, Odell DD, et al. Parenchymal preserving anatomic resections result in less pulmonary function loss in patients with Stage I non-small cell lung cancer. J Cardiothorac Surg 2015;10:49.
- 95. Kobayashi N, Kobayashi K, Kikuchi S, et al. Long-term pulmonary function after surgery for lung cancer. Interact Cardiovasc Thorac Surg 2017;24:727-32.
- 96. Helminen O, Valo J, Andersen H, et al. Thoracoscopic segmentectomy with simple routine bronchoscopic inflation for intersegmental plane identification: short and mid-term outcomes compared with lobectomy. J Thorac Dis 2020;12:3073-84.
- 97. Nagamatsu Y, Maeshiro K, Kimura NY, et al. Long-term recovery of exercise capacity and pulmonary function after lobectomy. J Thorac Cardiovasc Surg 2007;134:1273-8.
- Nakata M, Saeki H, Yokoyama N, et al. Pulmonary function after lobectomy: video-assisted thoracic surgery versus thoracotomy. Ann Thorac Surg 2000;70:938-41.
- Charloux A, Quoix E. Lung segmentectomy: does it offer a real functional benefit over lobectomy?. Eur Respir Rev 2017;26:170079.
- 100. Kim SJ, Lee YJ, Park JS, et al. Changes in pulmonary function in lung cancer patients after video-assisted thoracic surgery. Ann Thorac Surg 2015;99:210-7.
- 101. Suzuki K, Watanabe SI, Wakabayashi M, et al. A singlearm study of sublobar resection for ground-glass opacity dominant peripheral lung cancer. J Thorac Cardiovasc Surg 2022;163:289-301.e2.
- 102. Février E, Yip R, Becker BJ, et al. Change in quality of life of stage IA lung cancer patients after sublobar resection

and lobectomy. J Thorac Dis 2020;12:3488-99.

- 103. Handy JR Jr, Asaph JW, Douville EC, et al. Does videoassisted thoracoscopic lobectomy for lung cancer provide improved functional outcomes compared with open lobectomy? Eur J Cardiothorac Surg 2010;37:451-5.
- 104. Rizk NP, Ghanie A, Hsu M, et al. A prospective trial comparing pain and quality of life measures after anatomic lung resection using thoracoscopy or thoracotomy. Ann Thorac Surg 2014;98:1160-6.
- 105.Khullar OV, Rajaei MH, Force SD, et al. Pilot Study to Integrate Patient Reported Outcomes After Lung Cancer Operations Into The Society of Thoracic Surgeons Database. Ann Thorac Surg 2017;104:245-53.
- 106. Anami K, Horie J, Hirayama Y, et al. Changes in exercise tolerance and quality of life are unrelated in lung cancer survivors who undergo video-assisted thoracic surgery. J Phys Ther Sci 2018;30:467-73.
- 107.Fernando HC, Landreneau RJ, Mandrekar SJ, et al. Analysis of longitudinal quality-of-life data in highrisk operable patients with lung cancer: results from the ACOSOG Z4032 (Alliance) multicenter randomized trial. J Thorac Cardiovasc Surg 2015;149:718-25; discussion 725-6.
- 108. Schwartz RM, Yip R, Olkin I, et al. Impact of surgery for stage IA non-small-cell lung cancer on patient quality of life. J Community Support Oncol 2016;14:37-44.
- 109. Schwartz RM, Alpert N, Rosenzweig K, et al. Changes in quality of life after surgery or radiotherapy in early-stage lung cancer. J Thorac Dis 2019;11:154-61.
- 110. Sarna L, Cooley ME, Brown JK, et al. Women with lung cancer: quality of life after thoracotomy: a 6-month prospective study. Cancer Nurs 2010;33:85-92.
- 111. Schwartz RM, Yip R, Flores RM, et al. The impact of resection method and patient factors on quality of life among stage IA non-small cell lung cancer surgical patients. J Surg Oncol 2017;115:173-80.
- 112. Möller A, Sartipy U. Long-term health-related quality of life following surgery for lung cancer. Eur J Cardiothorac Surg 2012;41:362-7.
- 113.Möller A, Sartipy U. Changes in quality of life after lung surgery in old and young patients: are they similar? World J Surg 2010;34:684-91.
- 114.Pompili C, Brunelli A, Refai M, et al. Does chronic obstructive pulmonary disease affect postoperative quality of life in patients undergoing lobectomy for lung cancer? A case-matched study. Eur J Cardiothorac Surg 2010;37:525-30.
- 115. Salati M, Brunelli A, Xiumè F, et al. Quality of life in the

2382

Journal of Thoracic Disease, Vol 14, No 6 June 2022

elderly after major lung resection for lung cancer. Interact Cardiovasc Thorac Surg 2009;8:79-83.

- 116.Brunelli A, Socci L, Refai M, et al. Quality of life before and after major lung resection for lung cancer: a prospective follow-up analysis. Ann Thorac Surg 2007;84:410-6.
- 117.Handy JR Jr, Asaph JW, Skokan L, et al. What happens to patients undergoing lung cancer surgery? Outcomes and quality of life before and after surgery. Chest 2002;122:21-30.
- 118.Xu GW, Xie MR, Wu HR, et al. A prospective study examining the impact of uniportal video-assisted thoracic surgery on the short-term quality of life in patients with lung cancer. Thorac Cancer 2020;11:612-8.
- 119.Pompili C, Koller M, Velikova G, et al. EORTC QLQ-C30 summary score reliably detects changes in QoL three months after anatomic lung resection for Non-Small Cell Lung Cancer (NSCLC). Lung Cancer 2018;123:149-54.
- 120.Nugent SM, Golden SE, Hooker ER, et al. Longitudinal Health-related Quality of Life among Individuals Considering Treatment for Stage I Non-Small-Cell Lung Cancer. Ann Am Thorac Soc 2020;17:988-97.
- 121. Avery KNL, Blazeby JM, Chalmers KA, et al. Impact on Health-Related Quality of Life of Video-Assisted Thoracoscopic Surgery for Lung Cancer. Ann Surg Oncol 2020;27:1259-71.
- 122.Burfeind WR Jr, Tong BC, O'Branski E, et al. Quality of life outcomes are equivalent after lobectomy in the elderly. J Thorac Cardiovasc Surg 2008;136:597-604.
- 123.Balduyck B, Hendriks J, Lauwers P, et al. Quality of life evolution after lung cancer surgery: a prospective study in 100 patients. Lung Cancer 2007;56:423-31.
- 124. Alberts L, Wolff HB, Kastelijn EA, et al. Patient-reported Outcomes After the Treatment of Early Stage Non-smallcell Lung Cancer With Stereotactic Body Radiotherapy Compared With Surgery. Clin Lung Cancer 2019;20:370-377.e3.
- 125. Balduyck B, Hendriks J, Lauwers P, et al. Quality of life evolution after lung cancer surgery in septuagenarians: a prospective study. Eur J Cardiothorac Surg 2009;35:1070-5; discussion 1075.
- 126. Schulte T, Schniewind B, Dohrmann P, et al. The extent of lung parenchyma resection significantly impacts long-term quality of life in patients with non-small cell lung cancer. Chest 2009;135:322-9.
- 127. Schulte T, Schniewind B, Walter J, et al. Age-related impairment of quality of life after lung resection for non-

small cell lung cancer. Lung Cancer 2010;68:115-20.

- 128.Ilonen IK, Räsänen JV, Knuuttila A, et al. Quality of life following lobectomy or bilobectomy for non-small cell lung cancer, a two-year prospective follow-up study. Lung Cancer 2010;70:347-51.
- 129.Kenny PM, King MT, Viney RC, et al. Quality of life and survival in the 2 years after surgery for non small-cell lung cancer. J Clin Oncol 2008;26:233-41.
- 130. Balduyck B, Hendriks J, Lauwers P, et al. Quality of life after lung cancer surgery: a prospective pilot study comparing bronchial sleeve lobectomy with pneumonectomy. J Thorac Oncol 2008;3:604-8.
- 131. Wyrwich KW, Fihn SD, Tierney WM, et al. Clinically important changes in health-related quality of life for patients with chronic obstructive pulmonary disease: an expert consensus panel report. J Gen Intern Med 2003;18:196-202.
- 132. Wyrwich KW, Metz SM, Kroenke K, et al. Measuring patient and clinician perspectives to evaluate change in health-related quality of life among patients with chronic obstructive pulmonary disease. J Gen Intern Med 2007;22:161-70.
- 133.Maringwa JT, Quinten C, King M, et al. Minimal important differences for interpreting health-related quality of life scores from the EORTC QLQ-C30 in lung cancer patients participating in randomized controlled trials. Support Care Cancer 2011;19:1753-60.
- 134. Yost KJ, Eton DT, Garcia SF, et al. Minimally important differences were estimated for six Patient-Reported Outcomes Measurement Information System-Cancer scales in advanced-stage cancer patients. J Clin Epidemiol 2011;64:507-16.
- 135.Osoba D, Rodrigues G, Myles J, et al. Interpreting the significance of changes in health-related quality-of-life scores. J Clin Oncol 1998;16:139-44.
- 136. Sloan JA. Assessing the minimally clinically significant difference: scientific considerations, challenges and solutions. COPD 2005;2:57-62.
- 137.Möller A, Sartipy U. Predictors of postoperative quality of life after surgery for lung cancer. J Thorac Oncol 2012;7:406-11.
- 138. Balduyck B, Sardari Nia P, Cogen A, et al. The effect of smoking cessation on quality of life after lung cancer surgery. Eur J Cardiothorac Surg 2011;40:1432-7; discussion 1437-8.
- 139.Pompili C, Brunelli A, Xiumé F, et al. Predictors of postoperative decline in quality of life after major lung resections. Eur J Cardiothorac Surg 2011;39:732-7.

Detterbeck et al. Evidence for resection extent in healthy patients

- 140. Sartipy U. Influence of gender on quality of life after lung surgery. Eur J Cardiothorac Surg 2010;37:802-6.
- 141. Landreneau RJ, Mack MJ, Hazelrigg SR, et al. Prevalence of chronic pain after pulmonary resection by thoracotomy or video-assisted thoracic surgery. J Thorac Cardiovasc Surg 1994;107:1079-85; discussion 1085-6.
- 142. Maguire MF, Ravenscroft A, Beggs D, et al. A questionnaire study investigating the prevalence of the neuropathic component of chronic pain after thoracic surgery. Eur J Cardiothorac Surg 2006;29:800-5.
- 143.Bayman EO, Brennan TJ. Incidence and severity of chronic pain at 3 and 6 months after thoracotomy: metaanalysis. J Pain 2014;15:887-97.
- 144.Kwon ST, Zhao L, Reddy RM, et al. Evaluation of acute and chronic pain outcomes after robotic, video-assisted thoracoscopic surgery, or open anatomic pulmonary resection. J Thorac Cardiovasc Surg 2017;154:652-659.e1.
- 145.Brown LM, Kratz A, Verba S, et al. Pain and Opioid Use After Thoracic Surgery: Where We Are and Where We Need To Go. Ann Thorac Surg 2020;109:1638-45.
- 146. Kiser AC, Detterbeck FC. General aspects of surgical treatment. In: Detterbeck FC, Rivera MP, Socinski MA, et al. editors. Diagnosis and Treatment of Lung Cancer: An Evidence-Based Guide for the Practicing Clinician. Philadelphia, PA: W.B. Saunders, 2001:133-47.
- 147. Howington JA, Blum MG, Chang AC, et al. Treatment of Stage I and II Non-small Cell Lung Cancer. Chest 2013;143:e278S-e313S.
- 148. National Comprehensive CN. NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines) Non-Small Cell Lung Cancer Version 7.2019 - August 30, 2019 2019.
- 149.Mohiuddin K, Haneuse S, Sofer T, et al. Relationship between margin distance and local recurrence among patients undergoing wedge resection for small (≤2 cm) non-small cell lung cancer. J Thorac Cardiovasc Surg 2014;147:1169-75; discussion 1175-7.
- 150. Maurizi G, D'Andrilli A, Ciccone AM, et al. Margin Distance Does Not Influence Recurrence and Survival After Wedge Resection for Lung Cancer. Ann Thorac Surg 2015;100:918-24; discussion 924-5.
- 151.Sienel W, Stremmel C, Kirschbaum A, et al. Frequency of local recurrence following segmentectomy of stage IA nonsmall cell lung cancer is influenced by segment localisation and width of resection margins--implications for patient selection for segmentectomy. Eur J Cardiothorac Surg 2007;31:522-7; discussion 527-8.
- 152.Moon Y, Lee KY, Moon SW, et al. Sublobar Resection Margin Width Does Not Affect Recurrence of Clinical

N0 Non-small Cell Lung Cancer Presenting as GGO-Predominant Nodule of 3 cm or Less. World J Surg 2017;41:472-9.

- 153.Dolan D, Swanson SJ, Gill R, et al. Survival and Recurrence Following Wedge Resection Versus Lobectomy for Early-Stage Non-Small Cell Lung Cancer. Semin Thorac Cardiovasc Surg 2022;34:712-23.
- 154.El-Sherif A, Fernando HC, Santos R, et al. Margin and local recurrence after sublobar resection of non-small cell lung cancer. Ann Surg Oncol 2007;14:2400-5.
- 155.Wolf AS, Swanson SJ, Yip R, et al. The Impact of Margins on Outcomes After Wedge Resection for Stage I Non-Small Cell Lung Cancer. Ann Thorac Surg 2017;104:1171-8.
- 156.Masai K, Sakurai H, Sukeda A, et al. Prognostic Impact of Margin Distance and Tumor Spread Through Air Spaces in Limited Resection for Primary Lung Cancer. J Thorac Oncol 2017;12:1788-97.
- 157. Sawabata N, Maeda H, Matsumura A, et al. Clinical implications of the margin cytology findings and margin/ tumor size ratio in patients who underwent pulmonary excision for peripheral non-small cell lung cancer. Semin Thorac Cardiovasc Surg 2022;34:712-23.
- 158. Takahashi N, Sawabata N, Kawamura M, et al. Optimal sublobar resection for c-stage I non-small cell lung cancer: significance of margin distance to tumor size ratio and margin cytology (Supplementary analysis of KLSG-0801): complete republication. Gen Thorac Cardiovasc Surg 2019;67:690-6.
- 159. Fernando HC, Landreneau RJ, Mandrekar SJ, et al. Impact of brachytherapy on local recurrence rates after sublobar resection: results from ACOSOG Z4032 (Alliance), a phase III randomized trial for high-risk operable nonsmall-cell lung cancer. J Clin Oncol 2014;32:2456-62.
- 160. Tamura M, Matsumoto I, Tanaka Y, et al. Comparison Between Stereotactic Radiotherapy and Sublobar Resection for Non-Small Cell Lung Cancer. Ann Thorac Surg 2019;107:1544-50.
- 161.Moon Y, Lee KY, Park JK. Margin Width of Resected Lepidic Lung Cancer Does Not Affect Recurrence After Sublobar Resection. World J Surg 2018;42:1449-57.
- 162. Moon Y, Park JK, Lee KY. The Effect of Resection Margin Distance and Invasive Component Size on Recurrence After Sublobar Resection in Patients With Small (≤2 Cm) Lung Adenocarcinoma. World J Surg 2020;44:990-7.
- 163.Schuchert MJ, Pettiford BL, Keeley S, et al. Anatomic segmentectomy in the treatment of stage I non-small cell lung cancer. Ann Thorac Surg 2007;84:926-32; discussion

932-3.

- 164. Sawabata N, Ohta M, Matsumura A, et al. Optimal distance of malignant negative margin in excision of nonsmall cell lung cancer: a multicenter prospective study. Ann Thorac Surg 2004;77:415-20.
- 165. Kent M, Landreneau R, Mandrekar S, et al. Segmentectomy versus wedge resection for non-small cell lung cancer in high-risk operable patients. Ann Thorac Surg 2013;96:1747-54; discussion 1754-5.
- 166. Ren Y, Xie H, Dai C, et al. Prognostic Impact of Tumor Spread Through Air Spaces in Sublobar Resection for 1A Lung Adenocarcinoma Patients. Ann Surg Oncol 2019;26:1901-8.
- 167.Kadota K, Nitadori JI, Sima CS, et al. Tumor Spread through Air Spaces is an Important Pattern of Invasion and Impacts the Frequency and Location of Recurrences after Limited Resection for Small Stage I Lung Adenocarcinomas. J Thorac Oncol 2015;10:806-14.
- 168. Yanagawa N, Shiono S, Endo M, et al. Tumor spread through air spaces is a useful predictor of recurrence and prognosis in stage I lung squamous cell carcinoma, but not in stage II and III. Lung Cancer 2018;120:14-21.
- 169.Jia M, Yu S, Gao H, et al. Spread Through Air Spaces (STAS) in Lung Cancer: A Multiple-Perspective and Update Review. Cancer Manag Res 2020;12:2743-52.
- 170. Shiono S, Endo M, Suzuki K, et al. Spread through air spaces affects survival and recurrence of patients with clinical stage IA non-small cell lung cancer after wedge resection. J Thorac Dis 2020;12:2247-60.
- 171. Vaghjiani RG, Takahashi Y, Eguchi T, et al. Tumor Spread Through Air Spaces Is a Predictor of Occult Lymph Node Metastasis in Clinical Stage IA Lung Adenocarcinoma. J Thorac Oncol 2020;15:792-802.
- 172. Kim SK, Kim TJ, Chung MJ, et al. Lung Adenocarcinoma: CT Features Associated with Spread through Air Spaces. Radiology 2018;289:831-40.
- 173. Uruga H, Fujii T, Fujimori S, et al. Semiquantitative Assessment of Tumor Spread through Air Spaces (STAS) in Early-Stage Lung Adenocarcinomas. J Thorac Oncol 2017;12:1046-51.
- 174. Han YB, Kim H, Mino-Kenudson M, et al. Tumor spread through air spaces (STAS): prognostic significance of grading in non-small cell lung cancer. Mod Pathol 2021;34:549-61.
- 175. Toyokawa G, Yamada Y, Tagawa T, et al. Computed tomography features of resected lung adenocarcinomas with spread through air spaces. J Thorac Cardiovasc Surg 2018;156:1670-1676.e4.

- 176. Chen D, Mao Y, Wen J, et al. Tumor Spread Through Air Spaces in Non-Small Cell Lung Cancer: A Systematic Review and Meta-Analysis. Ann Thorac Surg 2019;108:945-54.
- 177.Dai C, Xie H, Su H, et al. Tumor Spread through Air Spaces Affects the Recurrence and Overall Survival in Patients with Lung Adenocarcinoma >2 to 3 cm. J Thorac Oncol 2017;12:1052-60.
- 178.Kadota K, Kushida Y, Kagawa S, et al. Limited Resection Is Associated With a Higher Risk of Locoregional Recurrence than Lobectomy in Stage I Lung Adenocarcinoma With Tumor Spread Through Air Spaces. Am J Surg Pathol 2019;43:1033-41.
- 179. Song T, Jiang L, Zhuo Z, et al. Impacts of thoracoscopic surgery and high grade histologic subtypes on spread through air spaces in small stage I lung adenocarcinomas. J Cancer Res Clin Oncol 2019;145:2375-82.
- 180.Hu SY, Hsieh MS, Hsu HH, et al. Correlation of tumor spread through air spaces and clinicopathological characteristics in surgically resected lung adenocarcinomas. Lung Cancer 2018;126:189-93.
- 181. Chae M, Jeon JH, Chung JH, et al. Prognostic significance of tumor spread through air spaces in patients with stage IA part-solid lung adenocarcinoma after sublobar resection. Lung Cancer 2021;152:21-6.
- 182. Shiono S, Endo M, Suzuki K, et al. Spread Through Air Spaces Is a Prognostic Factor in Sublobar Resection of Non-Small Cell Lung Cancer. Ann Thorac Surg 2018;106:354-60.
- 183. Toyokawa G, Yamada Y, Tagawa T, et al. Significance of Spread Through Air Spaces in Resected Pathological Stage I Lung Adenocarcinoma. Ann Thorac Surg 2018;105:1655-63.
- 184.Kadota K, Kushida Y, Katsuki N, et al. Tumor Spread Through Air Spaces Is an Independent Predictor of Recurrence-free Survival in Patients With Resected Lung Squamous Cell Carcinoma. Am J Surg Pathol 2017;41:1077-86.
- 185.Kagimoto A, Tsutani Y, Kushitani K, et al. Segmentectomy vs Lobectomy for Clinical Stage IA Lung Adenocarcinoma With Spread Through Air Spaces. Ann Thorac Surg 2021;112:935-43.
- 186. Toyokawa G, Yamada Y, Tagawa T, et al. Significance of spread through air spaces in early-stage lung adenocarcinomas undergoing limited resection. Thorac Cancer 2018;9:1255-61.
- 187. Valastyan S, Weinberg RA. Tumor metastasis: molecular insights and evolving paradigms. Cell 2011;147:275-92.

Detterbeck et al. Evidence for resection extent in healthy patients

- Wan L, Pantel K, Kang Y. Tumor metastasis: moving new biological insights into the clinic. Nat Med 2013;19:1450-64.
- 189. Weinberg RA. Mechanisms of malignant progression. Carcinogenesis 2008;29:1092-5.
- 190. Angelova M, Mlecnik B, Vasaturo A, et al. Evolution of Metastases in Space and Time under Immune Selection. Cell 2018;175:751-765.e16.
- 191. Nishio W, Yoshimura M, Maniwa Y, et al. Re-Assessment of Intentional Extended Segmentectomy for Clinical

Cite this article as: Detterbeck FC, Mase VJ Jr, Li AX, Kumbasar U, Bade BC, Park HS, Decker RH, Madoff DC, Woodard GA, Brandt WS, Blasberg JD. A guide for managing patients with stage I NSCLC: deciding between lobectomy, segmentectomy, wedge, SBRT and ablation—part 2: systematic review of evidence regarding resection extent in generally healthy patients. J Thorac Dis 2022;14(6):2357-2386. doi: 10.21037/jtd-21-1824 T1aN0 Non-Small Cell Lung Cancer. Ann Thorac Surg 2016;102:1702-10.

- 192. Roman M, Labbouz S, Valtzoglou V, et al. Lobectomy vs. segmentectomy. A propensity score matched comparison of outcomes. Eur J Surg Oncol 2019;45:845-50.
- 193. Ueda K, Tanaka T, Hayashi M, et al. What proportion of lung cancers can be operated by segmentectomy? A computed-tomography-based simulation. Eur J Cardiothorac Surg 2012;41:341-5.

2386

Supplementary file (Part 2 paper)

Table of Contents	Page
Tables	
Table S2-1: Adjusted perioperative morbidity and mortality studies	2
Table S2-2: Summary of evidence in generally healthy patients with typical (i.e., solid) tumors	3
Figures	
Figure S2-1: Graphic depiction of outcomes in Table 1: segmentectomy vs. lobectomy	4
Figure S2-2: Graphic depiction of outcomes in Table 2: wedge/sublobar resection vs. lobectomy	5
Figure S2-3: Graphic depiction of outcomes in Table 3: wedge vs. segmentectomy	6
Appendix	
Appendix 2-1: Tools to assess confidence in attribution to interventions	7-9
References	10-11

Table S2-1 Adjusted perioperative morbidity and mortality studiesOrdered by confidence that results reflect the effect of the treatment, resection extent

	1 st Author, year		St	udy ch	Study characteristics	stics			eriod ^b	Peric	Perioperative mortality	e morta	lity	G	Gr ≥3 Morbidity	orbidity	
NUMBE N Tag and the implication Value Vs. Continuents CH Equivable HS Equivable HS Continuents HS Contin HS	(reterence)	00000	Z	~~/	0+0.00 a	Lobe			d əw	Adjus	ted %	Adjuste	ad HR	Adjuste	% pe	Adjust∈	ad HR
STS $7,466^{\circ}$ $09-11$ $c-IIIA$ WVMNM30 1.2° 1.9° 2.4° 9° 3.5° 6° 3.5° 6.5° 3.5° 6.7° 2.5° 6.7° 2.5° 6.7° 6.7° 2.5° 6.7°		source	z	×1×	olage	VS.				Seg/W	Lobe	HR	٩	Seg/W	Lobe	HR	٩
STS5.288°09-11cl-IIAWVATSVH300.8 1.0° \sim \sim 3.5 6.7° 5.7° \sim STS2.004°09-11cl-IIAWOpenVH30 1.8 2.9° \sim \sim 6.8 12.4° \sim STS1,972°09-11cl-IIAWPetri < 60%	Linden 2014 (1)	STS	7,466 °		cl-IIIA	N		ΗΛ	30	1.2	1.9 ^d	1		4.5	_р 6	1	ı
STS2,00409-11cl-IIAWOpenVH30 1.82.9 $ -$ 6.812.4 $ -$ STS1,87209-11cl-IIAWFEV1<60%	Linden 2014 (1)	STS	5,288 °	09-11	cl-IIIA	≥	VATS	H	30	0.8	1.0 ^d	I	1	3.5	6.7 ^d	I	I
STS1,872°09-11cl·llaWFEV1<60%VH301.4 2.9^{d} $ 6.3$ 13.1^{d} $-$ STS1,068°09-11cl·llaWAge>800VH301.7 2.3^{d} $ 6.3$ 13.1^{d} $-$ NCDB75,11404-13cl·llaSLAge>800VH30 1.7 2.3^{d} $ 5.4$ 8.4^{d} $-$ Chinax10244°14-17MIxSegVATSM $ 0.8$ 1.6 $ -$	Linden 2014 (1)	STS	2,004 °		cl-IIIA	≥	Open	ΗΛ	30	1.8	2.9 ^d	I	1		12.4 ^d	I	ı
STS 1,068° 09-11 C-IIIA W Age=80 VH 30 1.7 2.3^{d} $ 5.4$ 8.4^{d} $-$ NCDB 75,114 04-13 cl-IIA SL $ H$ 90 3.3^{e} 3.5^{e} 0.87 $ -$ <td>Linden 2014 (1)</td> <td>STS</td> <td>1,872 °</td> <td>09-11</td> <td>cl-IIIA</td> <td>≥</td> <td>FEV1 <60%</td> <td>H</td> <td>30</td> <td>1.4</td> <td>2.9^d</td> <td>I</td> <td>1</td> <td></td> <td>13.1 ^d</td> <td>I</td> <td>I</td>	Linden 2014 (1)	STS	1,872 °	09-11	cl-IIIA	≥	FEV1 <60%	H	30	1.4	2.9 ^d	I	1		13.1 ^d	I	I
NCDB75,114 $0.4-13$ $cl-IA$ SLMCDR 0.6 3.3° 3.5° 0.87 $ -$ <t< td=""><td>Linden 2014 (1)</td><td>STS</td><td>1,068 °</td><td></td><td>cl-IIIA</td><td>≥</td><td>Age ≥80</td><td>ΗΛ</td><td>30</td><td>1.7</td><td>2.3 ^d</td><td>I</td><td>1</td><td>5.4</td><td>8.4 ^d</td><td>I</td><td>ı</td></t<>	Linden 2014 (1)	STS	1,068 °		cl-IIIA	≥	Age ≥80	ΗΛ	30	1.7	2.3 ^d	I	1	5.4	8.4 ^d	I	ı
China x10 244 ^e 14-17 Ci-liA SL Age=565, VATS H - 0.8 1.6 <1 NS - - - - - - - - - - - - - - - - - 1.6 <1 NS = - - - 0.8 1.6 1.6 NS - - - - - - - - - - - - - - - - - - - 0.8 1.6 0.31 - - 0 0 0 0 0 0 0 1.3 1 - - - 0 <td>Stokes 2018 (2)</td> <td>NCDB</td> <td>75,114</td> <td>04-13</td> <td>cl-IIA</td> <td>SL</td> <td></td> <td>т</td> <td>06</td> <td>3.3 °</td> <td>3.5 ^e</td> <td>0.87</td> <td>ı</td> <td>1</td> <td>ı</td> <td>1</td> <td>ı</td>	Stokes 2018 (2)	NCDB	75,114	04-13	cl-IIA	SL		т	06	3.3 °	3.5 ^e	0.87	ı	1	ı	1	ı
Swiss x2 690 14-17 Mix Seg VATS M - 0.3° $ -$ <	Zhang 2019 (3)	China x10	244 °	14-17		SL	Age ≥65, VATS	т	1	0.8	1.6	$\overline{\nabla}$	NS	I	I	I	I
NCDB $20,944$ $03-11$ $c A1,2$ Seg M 30 1.6° 1.6° NS $ -$	Bedat 2019 (4)	Swiss x2	069	14-17		Seg	VATS	Σ	1	0.8 °	0.4 ^e	ı	1			0.93 ^f	NS
NCDB $27,015$ $03-11$ $[A1,2]$ W M 30 1.5° 1.6° 0.05 $ -$	Khullar 2015 (5)	NCDB	20,944	03-11		Seg		Σ	8	1.6 ^e	1.6 ^e	0.87	SN	I	ı	Т	I
Japan x1 205 07-15 cl-IIA SL Age≥75 M - 0 ^e 0.9 ^e - - 5 10 0.29 NCDB 71,171 03-11 cl-IIIA SL L 30 2 ^{eg} 2.2 ^e 0.9 NS -	Khullar 2015 (5)	NCDB		03-11	cIA1,2	≥		Σ	30	1.5 [°]	1.6 ^e	0.72	.005	ı	•	I	ı
NCDB 71,171 03-11 cl-IIIA SL L 30 2 ^{e,g} 2.2 ^e 0.9 NS -	Tsutani 2018 (6)	-	205	07-15	cl-IIA	SL	Age ≥75	Μ		0 ^e	° 0.0	T	-	5	10	0.29	.048
NCDB 19,083 03-11 cl-IIIA SL Age≥75 L 30 0.76 .005	Husain 2015 (7)		71,171	03-11		SL		_	30	2 ^{e,g}	2.2 ^e	0.9	SN	1	ı	I	ı
	Husain 2015 (7)	NCDB	19,083	03-11	cl-IIIA	SL	Age ≥75	_	30	ı	I	0.76	.005	ı	ı	ı	ı

or wedge resection vs lobectomy, 2000–21, ≥50 patients per arm. Reference is lobectomy (HR <1 means lower morbidity/mortality for segment/wedge); Light green shading highlights statistically significant difference (lighter shade = univariable; darker = multivariable); Bold inclusion criteria: studies using multivariable of propensity adjustment to compare perioperative morbiolity/mortality after segmentectomy nighlights differences that are somewhat clinically meaningful (see definition in Part 1 paper).

8th edition stage classification (reported stage is translated into current 8th edition nomenclature for the sake of uniformity and contemporary application); ^b, time period for assessment of Morbidity and Mortality (days); ^c, propensity matched pairs (total); ^d, includes obectomy and segmentectomy; ^e, Unadjusted data; ^f, cardiopulmonary complications (any grade); ^g, data for wedge resection

Confid RE tmt effect, Confidence that results reflect the effect of the treatment (extent of resection) vs. confounding factors; FEV1, forced expiratory volume in 1 second; H, high confidence; HR, hazard ratio; L, low confidence; Lobe, lobectomy; M, moderate confidence; Mix, mixture of a variety of diagnoses (NSCLC, metastases, benign); NCDB, US national cancer database; NS, not statistically significant; Seg, segmentectomy; SL, sublobar resection (segmentectomy or wedge); VATS, video-assisted thoracic surgery; VH, very high confidence; W, wedge; Yrs, years (of patient accrual).

		ment _obe)		Weo (vs. L			dge gment)
	Effect	Conf	Eff	ect	Conf	Effect	Conf
Short-term (90-da	ay) outcoi	mes					
Mortality	=	++++	=	=	+++	=	+
Morbidity	=	+++	=	-	+++	=	+
QOL 30-day	= ^a	0	=	а	0	-	-
QOL 90-day	= ^a	0	=	а	0	-	-
Pain VATS	= ^a	0	=	а	0	-	-
Pain open	= ^a	0	=	а	0	-	-
Intermediate (1-2	year) out	comes					
Δ FEV1	=	++	=	⁄↑	0	-	-
Dyspnea	=/↑ ^a	0		↑ ^a	0	-	-
QOL VATS	= ^a	0	=	а	0	-	-
Pain VATS	= ^a	0	=	а	0	-	-
QOL open	= ^a	0	=	а	0	-	-
Pain open	= ^a	0	=	а	0	-	-
Long-term (5-yea	r) outcom	nes					
OS	Ļ	+	↓	↓ _	++	\downarrow	+
LCSS	Ļ	+			+	\downarrow	+
FFR	=/↓ ^a	0	=/	↓ ^a	0	-	-
LR- FFR	=/↓ ^a	0	=/	↓ ^a	0	-	-

Table S2-2 Summary of evidence in generally healthy patients with typical (i.e., solid) tumors

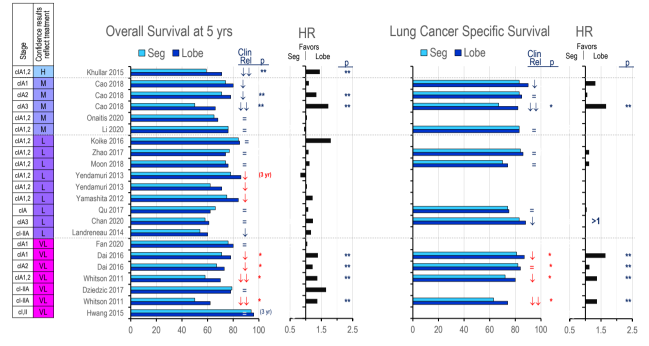
Qualitative assessment of the impact of treatment approaches on various key outcome measures and the confidence in the evidence. Differences are categorized by degree of clinically meaningful differences as defined in the legend insert. The reference (for improvement or worsening) is the treatment in parentheses.

	Effect	Confid	ence in / con-
$\uparrow\uparrow\uparrow$	2x meaningful	sistend	cy of evidence
	improvement		
$\uparrow\uparrow$	Meaningful improvement	++++	Very High
Î	Somewhat better	+++	High
=	Similar	++	Moderate
Ļ	Somewhat worse	+	Low
$\downarrow\downarrow$	Meaningful worsening	0	Very Low
$\downarrow\downarrow\downarrow\downarrow$	2x meaningful worsening	Extpol	Extrapolation

A clinically "meaningful" difference is defined as \geq 10-unit difference, with "somewhat" being half of the meaningful difference. The units of measure (for categories in parentheses) are: normalized scale points (QOL); 5-year actuarial rate (OS, LCSS); actuarial rate or simple incidence (recurrence, FFR); incidence of Gr \geq 3 treatment related complications (morbidity); absolute change in % FEV1 (PFTs in compromised patients). Different thresholds of "meaningful" are: 90-day mortality (2% difference); PFTs in healthy patients (20% difference in FEV1%).

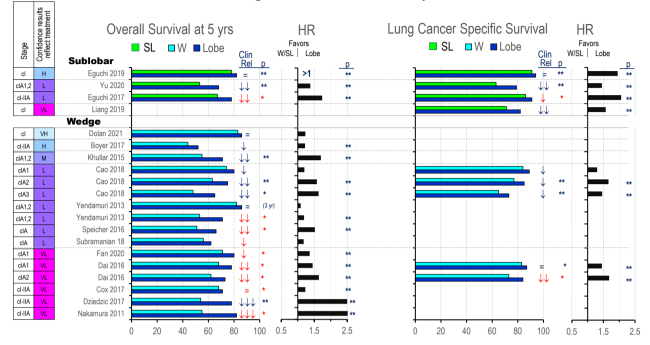
^a data for sublobar resection not parsed out to segment or wedge.

 Δ FEV1, change in FEV1 \geq 6 months; Conf, confidence in the evidence; Extpol, extrapolation (indirect evidence); FFR, freedom from recurrence (only recurrence counts as an event); Gr, grade; HR, hazard ratio; LCSS, lung cancer specific survival (only death due to lung cancer counts as an event); Lobe, lobectomy; LR-FFR, locoregional freedom from recurrence; OS, overall survival; PFT, pulmonary function tests; QOL, quality of life; VATS, video-assisted thoracic surgery.



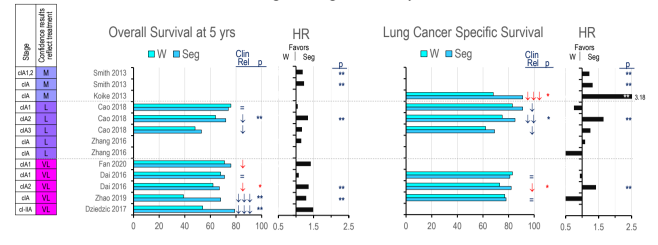
Segmentectomy vs. Lobectomy

Figure S2-1 Graphic depiction of outcomes in Table 1: segmentectomy vs. lobectomy.



Wedge/Sublobar vs. Lobectomy

Figure S2-2 Graphic depiction of outcomes in Table 2: wedge/sublobar resection vs. lobectomy.



Wedge vs. Segmentectomy

Figure S2-3 Graphic depiction of outcomes in Table 3: wedge vs. segmentectomy.

Legend (Figures S2-1,S2-2,S2-3): Graphic depiction of outcomes in *Tables 1-3*. Figure rows correspond to the respective table rows. Also depicted is the confidence that the outcomes reflect the treatment (*vs.* confounders), the level of clinical relevance and statistical significance.

	e results reflect reatment		Relevance of effect
VH	Very High	$\uparrow\uparrow\uparrow$	2x meaningfully better
н	High	$\uparrow\uparrow$	Meaningfully better
М	Moderate	1	Somewhat better
L	Low	=	Similar
VL	Very Low	\downarrow	Somewhat worse
See Tabl	e 1 for details	$\downarrow\downarrow$	Meaningfully worse
		$\downarrow\downarrow\downarrow\downarrow$	2x meaningfully worse

The HR reference is the larger resection, i.e., HR >1 reflects worse outcome compared with lobectomy (or segmentectomy in Figure S2-3).

Red font indicates unadjusted survival rates.

* reported as statistically significant by univariable analysis; ** reported as statistically significant by multivariable analysis; Clin Rel, clinical relevance of effect. A clinically relevant difference is defined as ≥5-point difference in the 5-year actuarial rate (overall survival, lung cancer specific survival). Details of this categorization is provided in the Part 1 paper (*Tab. S1-1*) (8). HR, hazard ratio; Lobe, lobectomy; Seg, segment; SL, sublobar resection; W, wedge; yrs, years.

Appendix 2-1: Tools to assess confidence in cause and effect attribution to the interventions in question

Assessment for confounding

ROBINS-I assessment

The Risk of Bias in Non-randomized Studies of Interventions (ROBINS-I) tool was used to assess included studies (9). This validated tool has gained acceptance for observational studies. The process involves identification of domains of bias for particular interventions, assessment of each study for potential bias relative to confounders and co-interventions in each domain, and aggregation of individual assessments into an overall risk of bias across studies. Studies are categorized as "low risk" if comparable to a well-done RCT, "moderate" is sound for a NRC but not comparable to a RCT, "serious" if at least one domain is not measured or controlled, and "critical risk" if internal or external data suggests residual confounding. It is suggested that critical studies be excluded from any systematic review (9).

In application of this tool, we found few that were low risk (2%), some that were moderate (18%); most were either serious (34%) or critical risk (45%). This illustrates problematic aspects of the ROBINS-I tool for our purpose. It is a generic tool designed largely to eliminate weak evidence. However, clinical care seeks to glean whatever information can be found; valuation rather than elimination seems more conducive to gaining an understanding of the strengths and pitfalls of the full scope of evidence. Furthermore, assessing the full spectrum of adjusted NRCs promotes uncovering reasons for discrepant results and nuances of which patients, tumors, and settings provide more convincing signs of efficacy.

Adapted assessment tool specific for this project

We adapted the ROBINS-I approach to the specific nature of our project. We identified 7 domains of potential confounding (detailed below) for the major long-term outcomes. We adopted a detailed approach that allows exploration of specific areas of confounding or patient and study characteristics. We adapted the rating of confounding, shifting from eliminating studies with potential confounding to assessing the impact of confounding on attribution of outcomes to the intervention of interest. This recognizes that the impact of unaddressed confounders can sometimes be ameliorated by the setting and study characteristics.

Domains of potential confounding Non-medical patient-related factors

Non-medical patient factors include age, sex, race, marital status, education level and income level. These factors have all been associated with long term outcomes in lung cancer patients (10,11). They can be thought of as influencing how aggressively patients want to be treated. Examples of factors that can affect the impact of such confounding include age cohorts under consideration, facility location, study region/ country (i.e. that might create greater or lesser uniformity of the study cohorts).

Medical patient-related factors

Comorbidities are more common in patients diagnosed with lung cancer than in a general population of similar age (12); these can account for competing causes of death. Most often a general measure of comorbidities such as the Charlson score is available. Such composite measures don't differentiate specific comorbidities or their severity. Ideally, additional information is available (e.g. FEV1, Performance status [PS]). Co-morbidities should not impact LCSS, since only a death due to lung cancer is counted as an event. (Consistent effect for OS and LCSS argues against major comorbidity confounding for OS).

Stage accuracy

The method and thoroughness of stage assessment differs among the interventions in question (e.g., wedge resections are often Nx). Additionally, until recently the SEER database only recorded best stage (clinical for non-surgical interventions, pathologic for surgery). Mitigating factors for discrepancies in stage assessment include use of PET, invasive mediastinal staging, risk of node involvement according to tumor characteristics (size, GG component).

Study time span

Often outcome studies encompass many years. The impact of trends over time is complicated. The proportion of resections involving sublobar resection is increasing as is the use of SBRT and ablation (13-16). The use of VATS is increasing, as is PET (17,18). There is also a trend towards detecting smaller size lung cancers, and an increase in lung cancers with a ground glass component (14,19,20). All of these factors potentially confound interpretation of studies: changing nature of tumors, type of resection, type of surgeon/radiotherpist and facilities at which they are performed—all of which are associated with differences in long-term outcomes.

Examples of factors contributing to the impact include the duration of the time span, whether adjustment is dichotomized or more differentiated, whether PET was used consistently, interactions with facility characteristics, tumor characteristics (size, GG component) and whether these are accounted for.

Setting characteristics

Facility characteristics are associated with discrepancies in the use of treatment modalities. For example, wedge resection may be associated with both the lowest volume and the highest volume hospitals, non-thoracic surgeons and nonacademic hospitals (13,21), and regional discrepancy in the use of SBRT and ablation is well documented (15). There are likely interactions between the setting and characteristics like details of pre-treatment evaluation, how tumors are detected, timeliness of care. Mitigating factors include the nature of the data source, breadth of facilities in question.

Treatment quality

Different treatment approaches may be associated with differences that affect outcomes, for example margin extent, use of adjuvant therapy, discrepancy in technical treatment factors (e.g. VATS), conversion to lobectomy if margins or nodes are concerning. All of these can produce discrepancies in factors other than the treatment intervention itself that can affect outcomes.

Favorable tumors

It is likely that tumors deemed more favorable are selected for lesser interventions (e.g. mostly GG, low PET activity, slow growth). It is clear that CT screening as well as incidental detection leads to an increased proportion of biologically more indolent tumors (22-24). Tumors with a ground glass (GG) appearance have a better prognosis (25). The presence of even a small GG component is associated with better outcomes (26,27). Prognosis correlates with the size of the solid component, not the GG component (25,28-33).

Methods of multivariable adjustment

Research involving large databases can provide an assessment of effects of a treatment in the "real world." However, ascribing an observed difference in outcomes to an intervention of interest requires assuming that nothing else is different—regarding the patients, the setting, the measurement of the outcomes etc. Since this is almost always not true, adjustment is necessary to mitigate the effect of confounding. It has become common to use propensity score analysis to accomplish this. It is worth explicitly noting several principles of this method. First, it can only adjust for known and observed factors - unmeasured factors remain a problem (e.g. severity of a condition, assessment of frailty). Second, propensity score analysis requires the assumption that any factors not included in the adjustment are "ignorable"i.e., not associated with who will or will not receive the intervention in question (34). Indeed, derivation of the propensity score should include all factors that may be related to the outcomes and/or the treatment decision (but not those related to outcomes alone) (35). However, most outcomes studies of limited resection or SBRT have omitted adjustment for factors that are clearly related to the choice of treatment (e.g., sicker patients, favorable tumors, type of treatment facility, time period). Third, the ability of propensity scores to mitigate the effect of confounding is variable; it depends on which adjustment method is used, characteristics of the population (e.g., whether treated and control groups are markedly skewed, have a large amount of overlap or one is contained in the other, number of events) (35-37).

There are many ways of using the propensity score to adjust for confounding: the most common are (I) propensity adjustment (PA) that uses the propensity score as an additional variable in a multivariable model, (II) propensity matching (PM), involving creation of 2 subsets (treatment and control) in which each treatment patient is paired with a control patient with an equal (or nearly equal) propensity score, (III) stratification, usually into quintiles, of the entire study population (PQ), with assessment of the treatment effect in each, and 4) inverse propensity weighting (PW) in which treated patients that were less likely to be treated (and vice versa) are weighed more heavily, essentially creating an equalized pseudopopulation. Which method is best depends on many factors: e.g., PM is not ideal with small samples, PW does not perform well in skewed populations, and PQ in survival analyses, but this is an oversimplification (35-37).

Because details of the propensity score development and the type of analysis affect how well the process can mitigate confounding effects, it is beneficial to perform additional analyses (different methods of adjustment, age groups, tumor size categories). Such additional analyses do not adjust for unmeasured factors or prove that they are ignorable, but if the observed effect is consistent it provides a degree of increased confidence that it is related to the intervention in question; in contrast if it is inconsistent there should be significant caution in attributing the effect in any one group to the intervention of interest. While specific techniques can diminish some of the limitations of each method, the complexity underscores that propensity score adjustment does not guarantee that an observed effect is related to the intervention in question.

Finally, it is not clear that propensity analysis adjusts for confounders better than multivariable adjustment models (e.g., cox regression) (35,36). Multivariable regression models the relationship between multiple covariates and outcome. Because simultaneous adjustment for multiple confounders is complex, a substantial sample size is needed-it is generally accepted that about 10 events are required for each included covariate. Propensity scoring models the relationship between confounders and the treatment assignment, thus collapsing all confounders into a single propensity score. In theory, propensity techniques may have an advantage when the number of confounders is large and the number of events is small. However, analyses have not clearly demonstrated that propensity methods provide a more accurate estimate of treatment effect than multivariable methods (35,36).

Assessment process

Two individuals independently assessed each study using the adapted tool; differences were resolved by discussion or a third assessment. There was agreement in most cases or only minor differences regarding adjacent degrees of concern in individual domains. It was rare that resolution of discrepant evaluations changed the overall study rating. Results of the consensus assessment are shown in the relevant tables. Additionally, each study was assessed using the ROBINS-I tool. Our adapted rating was generally consistent with the ROBINS-I rating, although our scale allowed a more differentiated range (we avoided the threshold for a NRC of being comparable to a well-done RCT, and tried to understand critical confounders instead of a threshold of "one and you're out" approach).

Additional information

Further detail (individual rating results, reasons for ratings etc.) available if desired.

References

- Linden PA, D'Amico TA, Perry Y, Saha-Chaudhuri P, Sheng S, Kim S, et al. Quantifying the safety benefits of wedge resection: a society of thoracic surgery database propensity-matched analysis. Ann Thorac Surg. 2014;98(5):1705-11; discussion 11-2.
- Stokes WA, Bronsert MR, Meguid RA, Blum MG, Jones BL, Koshy M, et al. Post-Treatment Mortality After Surgery and Stereotactic Body Radiotherapy for Early-Stage Non-Small-Cell Lung Cancer. J Clin Oncol. 2018;36(7):642-51.
- Zhang Z, Feng H, Zhao H, Hu J, Liu L, Liu Y, et al. Sublobar resection is associated with better perioperative outcomes in elderly patients with clinical stage I non-small cell lung cancer: a multicenter retrospective cohort study. J Thorac Dis. 2019;11(5):1838-48.
- Bédat B, Abdelnour-Berchtold E, Perneger T, Licker MJ, Stefani A, Krull M, et al. Comparison of postoperative complications between segmentectomy and lobectomy by video-assisted thoracic surgery: a multicenter study. J Cardiothorac Surg. 2019;14(1):189.
- Khullar OV, Liu Y, Gillespie T, Higgins KA, Ramalingam S, Lipscomb J, et al. Survival after sublobar resection versus lobectomy for clinical stage IA lung cancer: An analysis from the National Cancer Data Base. J Thorac Oncol. 2015;10(11):1625-33.
- Tsutani Y, Tsubokawa N, Ito M, Misumi K, Hanaki H, Miyata Y, et al. Postoperative complications and prognosis after lobar resection versus sublobar resection in elderly patients with clinical Stage I non-small-cell lung cancer. Eur J Cardiothorac Surg. 2018;53(2):366-71.
- Husain ZA, Kim AW, Yu JB, Decker RH, Corso CD. Defining the High-Risk Population for Mortality After Resection of Early Stage NSCLC. Clin Lung Cancer. 2015;16(6):e183-7.
- Detterbeck F, Blasberg J, Woodard G, Decker R, Kumbasar U, Park H, et al. A Guide for Managing Patients with Stage I NSCLC: Deciding between Lobectomy, Segmentectomy, Wedge, SBRT and Ablation. Part 1: A Guide to Decision-Making. J Thor Dis. 2022.
- Sterne JA, Hernan MA, Reeves BC, Savovic J, Berkman ND, Viswanathan M, et al. ROBINS-I: a tool for assessing risk of bias in non-randomised studies of interventions. BMJ. 2016;355:i4919.
- Albano JD, Ward E, Jemal A, Anderson R, Cokkinides VE, Murray T, et al. Cancer mortality in the United States by education level and race. J Natl Cancer Inst. 2007;99(18):1384-94.

- Wong MCS, Lao XQ, Ho KF, Goggins WB, Tse SLA. Incidence and mortality of lung cancer: global trends and association with socioeconomic status. Sci Rep. 2017;7(1):14300.
- Cho H, Mariotto AB, Mann BS, Klabunde CN, Feuer EJ. Assessing non-cancer-related health status of US cancer patients: other-cause survival and comorbidity prevalence. American journal of epidemiology. 2013;178(3):339-49.
- Kim AW, Detterbeck FC, Boffa DJ, Decker RH, Soulos PR, Cramer LD, et al. Characteristics associated with the use of nonanatomic resections among Medicare patients undergoing resections of early-stage lung cancer. Ann Thorac Surg. 2012;94(3):895-901.
- McMurry TL, Shah PM, Samson P, Robinson CG, Kozower BD. Treatment of stage I non-small cell lung cancer: What's trending? J Thorac Cardiovasc Surg. 2017;154(3):1080-7.
- Uhlig J, Ludwig JM, Goldberg SB, Chiang A, Blasberg JD, Kim HS. Survival Rates after Thermal Ablation versus Stereotactic Radiation Therapy for Stage 1 Non-Small Cell Lung Cancer: A National Cancer Database Study. Radiology. 2018;289(3):862-70.
- Vest MT, Herrin J, Soulos PR, Decker RH, Tanoue L, Michaud G, et al. Use of new treatment modalities for non-small cell lung cancer care in the Medicare population. Chest. 2013;143(2):429-35.
- 17. Detterbeck FC. Maintaining Aim at a Moving Target. J Thor Oncol. 2011;6(3):417-22.
- Detterbeck F, Molins L. Video-assisted thoracic surgery and open chest surgery in lung cancer treatment: present and future. J Vis Surg. 2016;2:173.
- Koike T, Yamato Y, Asamura H, Tsuchiya R, Sohara Y, Eguchi K, et al. Improvements in Surgical Results for Lung Cancer from 1989 to 1999 in Japan. J Thorac Oncol. 2009;4(11):1364-9.
- 20. Okami J, Shintani Y, Okumura M, Ito H, Ohtsuka T, Toyooka S, et al. Demographics, Safety and Quality, and Prognostic Information in Both the Seventh and Eighth Editions of the TNM Classification in 18,973 Surgical Cases of the Japanese Joint Committee of Lung Cancer Registry Database in 2010. J Thorac Oncol. 2019;14(2):212-22.
- Camposilvan I, Akhtar-Danesh N, Schneider L, Finley CJ. The effect of surgeon volume on procedure selection in non-small cell lung cancer surgeries. J Thorac Cardiovasc Surg. 2015;150(3):507-12.
- 22. Detterbeck F, Gibson C. Turning Gray: The Natural History of Lung Cancer Over Time. J Thorac Oncol. 2008;3(7):781-92.

- 23. Schabath MB, Massion PP, Thompson ZJ, Eschrich SA, Balagurunathan Y, Goldof D, et al. Differences in Patient Outcomes of Prevalence, Interval, and Screen-Detected Lung Cancers in the CT Arm of the National Lung Screening Trial. PLoS One. 2016;11(8):e0159880.
- Quadrelli S, Lyons G, Colt H, Chimondeguy D, Buero A. Clinical characteristics and prognosis of incidentally detected lung cancers. Int J Surg Oncol. 2015;2015:287604.
- 25. Mase VJ, Jr., Detterbeck FC. Approach to the Subsolid Nodule. Clin Chest Med. 2020;41(1):99-113.
- 26. Hattori A, Hirayama S, Matsunaga T, Hayashi T, Takamochi K, Oh S, et al. Distinct Clinicopathologic Characteristics and Prognosis Based on the Presence of Ground Glass Opacity Component in Clinical Stage IA Lung Adenocarcinoma. J Thorac Oncol. 2019;14(2):265-75.
- Miyoshi T, Aokage K, Katsumata S, Tane K, Ishii G, Tsuboi M. Ground-Glass Opacity Is a Strong Prognosticator for Pathologic Stage IA Lung Adenocarcinoma. Ann Thorac Surg. 2019;108(1):249-55.
- 28. Maeyashiki T, Suzuki K, Hattori A, Matsunaga T, Takamochi K, Oh S. The size of consolidation on thinsection computed tomography is a better predictor of survival than the maximum tumour dimension in resectable lung cancer. Eur J Cardiothorac Surg. 2013;43(5):915-8.
- 29. Murakawa T, Konoeda C, Ito T, Inoue Y, Sano A, Nagayama K, et al. The ground glass opacity component can be eliminated from the T-factor assessment of lung adenocarcinoma. Eur J Cardiothorac Surg. 2013;43(5):925-32.
- 30. Sawabata N, Kanzaki R, Sakamoto T, Kusumoto H, Kimura T, Nojiri T, et al. Clinical predictor of pre- or minimally invasive pulmonary adenocarcinoma: possibility of sub-classification of clinical T1a. European Journal of

Cardio-Thoracic Surgery. 2014;45(2):256-61.

- 31. Tsutani Y, Miyata Y, Nakayama H, Okumura S, Adachi S, Yoshimura M, et al. Prognostic significance of using solid versus whole tumor size on high-resolution computed tomography for predicting pathologic malignant grade of tumors in clinical stage IA lung adenocarcinoma: A multicenter study. J Thorac Cardiovasc Surg. 2012;143(3):607-12.
- 32. Tsutani Y, Miyata Y, Mimae T, Kushitani K, Takeshima Y, Yoshimura M, et al. The prognostic role of pathologic invasive component size, excluding lepidic growth, in stage I lung adenocarcinoma. J Thorac Cardiovasc Surg. 2013;146(3):580-5.
- 33. Yanagawa N, Shiono S, Abiko M, Ogata S-y, Sato T, Tamura G. New IASLC/ATS/ERS Classification and Invasive Tumor Size are Predictive of Disease Recurrence in Stage I Lung Adenocarcinoma. J Thorac Oncol. 2013;8(5):612-8.
- 34. Freemantle N, Marston L, Walters K, Wood J, Reynolds MR, Petersen I. Making inferences on treatment effects from real world data: propensity scores, confounding by indication, and other perils for the unwary in observational research. BMJ. 2013;347:f6409.
- 35. Elze MC, Gregson J, Baber U, Williamson E, Sartori S, Mehran R, et al. Comparison of Propensity Score Methods and Covariate Adjustment: Evaluation in 4 Cardiovascular Studies. J Am Coll Cardiol. 2017;69(3):345-57.
- Hade EM, Lu B. Bias associated with using the estimated propensity score as a regression covariate. Stat Med. 2014;33(1):74-87.
- Benedetto U, Head SJ, Angelini GD, Blackstone EH. Statistical primer: propensity score matching and its alternatives. Eur J Cardiothorac Surg. 2018;53(6):1112-7.