The Influence of Lead-Related Venous Obstruction on the Complexity and Outcomes of Transvenous Lead Extraction

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Abstract: Background: Little is known about lead-related venous stenosis/occlusion (LRVSO), and the influence of LRVSO on the complexity and outcomes of transvenous lead extraction (TLE) is debated in the literature. Methods: We performed a retrospective analysis of venograms from 2909 patients who underwent TLE between 2008 and 2021 at a high-volume center. Results: Advanced LRVSO was more common in elderly men with a high Charlson comorbidity index. Procedure duration, extraction of superfluous leads, occurrence of any technical difficulty, lead-to-lead binding, fracture of the lead being extracted, need to use alternative approach and lasso catheters or metal sheaths were found to be associated with LRVSO. The presence of LRVSO had no impact on the number of major complications including TLE-related tricuspid valve damage. The achievement of complete procedural or clinical success did not depend on the presence of LRVSO. Long-term mortality, in contrast to periprocedural and short-term mortality, was significantly worse in the groups with LRSVO. Conclusions: LRVSO can be considered as an additional TLE-related risk factor. The effect of LRVSO on major complications including periprocedural mortality and on short-term mortality has not been established. However, LRVSO has been associated with poor long-term survival.

Keywords: lead-related venous obstruction; transvenous lead extraction; lead extraction complications; lead extraction complexity

1. Background

Permanent cardiac pacing remains the leading treatment for patients with various rhythm disorders, conduction disturbances and ventricular arrhythmias. In recent years, we have also observed an increase in the implantation of more complex devices used in the prevention of sudden cardiac death and in the treatment of severe heart failure. In spite of technological progress over the last decade, conventional pacemakers, implantable cardioverter-defibrillators and cardiac resynchronization therapy (PM/ICD/CRT) devices still have endocardial leads. However, after the beginning of the endocardial pacing era only
Several studies have investigated lead-related venous stenosis/occlusion (LRVSO) [1–20]. Various lead-related problems (infectious and non-infectious) are an inherent component of permanent endocardial pacing, and transvenous lead extraction (TLE) is considered an essential technique in lead management strategy [21–24]. TLE is a complex procedure that sometimes may lead to fatal complications such as venous or cardiac injury. Carrying out the procedure is often associated with technical problems and requires additional approaches and tools [22–25]. There are numerous reports on the estimation of the real risk of the TLE [26–29] but none of them considered LRVSO as a predictor of procedure difficulties. Among twenty reports on LRVSO [1–20], only three papers analyzed the occurrence of LRVSO before TLE [2,6,14], and only two considered the influence of LRVSO on procedure complexity providing at the same time conflicting results. Among 20 reports on LRVSO only two studies were carried out in populations over 200 patients [1,2], 10 in 100–150 participants [3–12] and the remaining eight studies in populations consisting of 30–89 patients [13–20]. In this study, a total of 2909 TLE procedures were preceded by venography and LRVSO was documented in 2138 venograms. Ipsilateral venography before TLE is an integral part of the procedure (in the absence of contraindications for contrast intake).

**Goal of the Study**

The aim of this study was to determine the incidence of varying degrees of LRVSO and to examine the influence of LRVSO on procedure difficulty, complexity, major complications related to TLE, procedure effectiveness as well as mid- and long-term mortality after TLE.

**2. Methods**

**2.1. Study Population**

A post-hoc analysis of clinical data from 2909 patients undergoing transvenous lead extraction (TLE) between June 2008 and March 2021 at a single high-volume center was performed. All information regarding the patients and the procedures were entered into the computer database on a current basis. Patients with medical contraindications for venography (contrast intake) were excluded from the study.

Table 1 summarizes the most important information regarding the study population.

**Table 1. Characteristics of the study group.**

| Characteristic                                      | Mean/Number | SD/% |
|-----------------------------------------------------|-------------|------|
| Patient age during TLE [years]                      | 66.90       | 13.99|
| Patient age at first implantation [years]           | 58.51       | 15.67|
| Sex (% of female patients)                          | 1147        | 39.43%|
| Etiology: IHD, MI                                  | 1676        | 57.61%|
| Etiology: cardiomyopathy, valvular heart disease    | 448         | 15.40%|
| Etiology: congenital, channelopathies, neurocardiogenic, post cardiac surgery | 784 | 26.95%|
| LVEF [%]                                            | 48.89       | 15.21|
| Renal failure (any)                                 | 607         | 20.87%|
| Previous sternotomy                                 | 435         | 14.95%|
| Charlson comorbidity index [number of points]       | 4.775       | 3.625|
| Systemic infection (with pocket infection or not)   | 599         | 20.59%|
| Local (pocket) infection                            | 253         | 8.70% |
| Lead failure (replacement)                          | 1505        | 57.74%|
Table 1. Cont.

| Change of pacing mode/upgrading, downgrading | 176 | 6.05% |
|---------------------------------------------|-----|-------|
| Other (abandoned lead/prevention of abandonment (AF, superfluous leads), threatening/potentially threatening lead (loops, free ending, left heart, LDTD), other (MRI indications, cancer, painful pocket, loss of indications for pacing/ICD), regaining venous access (symptomatic occlusion, SVC syndrome, lead replacement/upgrading) | 374 | 12.86% |
| System: pacemaker (any) | 2013 | 69.20% |
| System: ICD-V, ICD-D | 281 | 22.93% |
| System: CRT-D | 667 | 7.80% |
| Dwell time of the oldest lead per patient before TLE [months] | 101.5 | 75.57 |
| Cumulative lead dwell time before TLE [years] | 15.31 | 12.925 |
| Major complications: all | 61 | 2.10% |
| Major complications (with rescue cardiac surgery) | 35 | 1.20% |
| Major complications (without rescue cardiac surgery) | 7 | 1.20% |
| Minor complications | 174 | 5.98% |

AF—atrial fibrillation, CRT-D—implantable cardioverter-defibrillator with resynchronization function, ICD—implantable cardioverter-defibrillator, ICD-D—dual chamber implantable cardioverter-defibrillator, ICD-V—single chamber implantable cardioverter-defibrillator, IHD—ischemic heart disease, LDTD—lead-dependent tricuspid dysfunction, LVEF—left ventricular ejection fraction, MI—myocardial infarction, MRI—magnetic resonance imaging, SVC—superior vena cava, TLE—transvenous lead extraction.

The current study uses data from a high-volume center that performs more than 200 TLE per year.

The percentage of serious complications is relatively higher compared to other reports, however, in the presented center, the most difficult procedures in the country are performed.

The first line tools used in study center are conventional mechanical sheaths, powered rotational mechanical sheaths and other instruments are second-line tools. Excimer laser sheaths are not used.

2.2. Venography

Preoperative venography was performed in 2909 patients submitted for transvenous lead extraction between June 2008 and March 2021 at our high-volume center. A peripheral intravenous catheter was placed in the peripheral arm vein on the side (or both sides of the chest) to be examined. All patients received an injection of 20–40 mL high-quality contrast medium (350 mg iodine/mL) Iomeron 350 into the peripheral arm vein on the side of endocardial lead implantation. Venous blood flow in the upper arm, neck and chest was recorded by cine-angiography. All images were acquired in the anteroposterior view. The venograms were obtained in a single plane (anterior–posterior) and stored on CD-ROM discs. An experienced cardiologist and experienced (trained by an interventional radiologist) cardiac surgeon reviewed the venograms, and venous patency was graded on a five-degree scale from normal flow to complete occlusion. All venograms were obtained in the same manner. Venographic analysis: at baseline, the narrowest and widest points of the target vessel for lead placement were identified by visual inspection to obtain minimum and maximum venous diameters, and measurements from two to three individually calibrated frames were averaged to determine the final status of the vein as no stenosis, mild stenosis (<50% narrowing), moderate stenosis (50–80% narrowing), severe stenosis (≥80% narrowing) and complete occlusion of the axillary (AxV), subclavian (ScV), innominate (brachiocephalic) (AnV) veins and superior vena cava (SVC). In spite of contrast injection in the arm vein on the side of the endocardial lead, regional collateral blood vessels and venous collateral blood flow in the neck enabled evaluation of the
brachiocephalic vein on the opposite side of the chest. What is the significance of this classification of vessel narrowing in clinical practice? LVRSO was graded according to our own, arbitrarily estimated, criteria, which rely to the remaining effective vein lumen necessary for different electrodes/catheters safe passage.

Mild narrowing: possible insertion of a new/additional lead using standard introducers, central venous catheters, permanent catheters for hemodialysis and there is a chance that the arteriovenous (AV) fistula will work properly.

Moderate narrowing: probable insertion of a new lead but hydrophilic guide wires and longer introducers are necessary, possible insertion of central venous catheters (troubles possible), possible insertion of permanent catheters for hemodialysis and there is a small chance that the AV fistula will work properly.

Severe narrowing: impossible insertion of a new lead, hydrophilic guide wires and longer introducers might be helpful, insertion of central venous catheters may be risky, chances to pass a catheter for hemodialysis without venoplasty are very small and there is no chance that the AV fistula will work properly.

Complete occlusion: no chance to pass a hydrophilic guide wire; only lead extraction and regaining venous access enables the insertion of a new lead.

Reuse of occluded veins and technical aspects of lead extraction/replacement depend not only on maximal venous narrowing but also on the length of the narrowing (the number of the affected vessels, too).

2.3. Lead Extraction Procedure

Lead extraction procedures were defined according to the most recent guidelines on management of lead-related complications (HRS 2017 and EHRA 2018) [21–23]. Indications for TLE and type of periprocedural complications were defined according to the 2017 HRS Expert Consensus Statement on Cardiovascular Implantable Electronic Device Lead Management and Extraction [22].

All procedures were performed using non-powered mechanical systems such as polypropylene Byrd dilator sheaths (Cook® Medical, Leechburg, PA, USA), mainly via the implant vein. If technical difficulties arose, alternative venous approaches and/or additional tools such as Evolution (Cook® Medical, Leechburg, PA, USA), TightRail (Spectranetix, Colorado Springs, CO, USA), lassos, basket catheters were used. Excimer laser sheaths were not used.

All extraction procedures were performed following different organizational models spanning 25 years of experience. In the initial era of lead extraction, the procedures were performed in the electrophysiology laboratory using intravenous analgesia/sedation; then the recommended safety precautions were observed to perform more complex and risky procedures in the operating theater, and finally in the hybrid room under general anesthesia. The core extraction team has consisted of the same very experienced TLE operator and a dedicated cardiac surgeon with an experienced echocardiographist over the last six years.

2.4. TEE Monitoring during TLE

TTE, pre- and postoperative TEE were mandatory (excluding contraindications) from the very beginning. Continuous transesophageal echocardiographic (TEE) monitoring has been an important standard tool over the last six years [30–32]. TEE in our series was performed using Philips iE33 or GE Vivid S 70 machines equipped with X7-2t Live 3D or 6VT-D probes. All recordings were archived and consisted of pre-procedural examination, navigation of lead removal and post-procedural evaluation of the efficacy of the procedure with an assessment of possible complications [30–32]. The intra-procedural phase of TEE monitoring allowed visualization of pulling on the cardiac walls and invagination of the right ventricle during lead removal, followed by a drop in systolic blood pressure in response to this maneuver. Continuous monitoring made it possible to clarify the cause of blood pressure fall during TLE [30–32].
2.5. Statistical Analysis

The Shapiro–Wilk test showed that most continuous variables were normally distributed. For uniformity, all continuous variables are presented as the mean ± standard deviation. In the first step the Kruskal–Wallis ANOVA test was used to determine whether there were statistically significant differences between groups. Next, the variables achieving \( p < 0.1 \) were compared using the nonparametric Chi\(^2\) test with Yates correction (dichotomous data) or the unpaired Mann–Whitney U test (continuous data), as appropriate. Comparisons were made between Groups 1 and 2 vs. Groups 4 and 5. A \( p \)-value less than 0.05 was considered as statistically significant. In order to assess the effect of LRVSO on mortality, Kaplan–Meier survival curves were plotted, the course of which was assessed using the log rank test. Statistical analysis was performed with Statistica version 13.3 (TIBCO Software Inc., Palo Alto, CA, USA).

2.6. Approval of the Bioethics Committee

All patients gave their informed written consent to undergo TLE and use anonymous data from their medical records, approved by the Bioethics Committee at the Regional Chamber of Physicians in Lublin No. 288/2018/KB/VII. The study was carried out in accordance with the ethical standards of the 1964 Declaration of Helsinki.

3. Results

Patient Groups

For the purposes of analysis, the study population was divided into five groups according to venogram results, namely Group 1—no stenosis (499 patients), 2—mild stenosis (574 pts), 3—moderate stenosis (605 pts), 4—severe stenosis (581 pts) and 5—total occlusion (650 pts). Only maximal venous narrowing was considered as a criterion in patient selection.

Tables 2–4 summarize specific patient-, system- and procedure-related risk factors for procedure complexity, efficacy, complications and long-term mortality after TLE.

Analysis of the clinical factors demonstrated that lead-related stenosis/occlusion correlated with patient age during TLE, male gender and Charlson comorbidity index. Other patient-related risk factors for major complications, i.e., indications for CIED implantation, functional NYHA class III and IV, decreased LVEF, renal failure and previous sternotomy were not related to LRVSO (Table 2).

Analysis of CIED systems and history of pacing showed that venous stenosis or lead-related (LR) total venous occlusion were more frequent in CRT-D recipients. Patients with ICD (VVI, DDD) were less likely, albeit insignificantly to have total venous occlusion.

Patients with redundant loops of the lead before TLE, leads with proximal end in the coronary sinus vein (CSV) and a higher number of CIED-related procedures before lead extraction were more likely to have severe venous stenosis or total occlusion.

Patients with severe venous stenosis or LR total venous occlusion had more risk factors for major complications (MC) and higher procedure complexity estimated with the SAFeTY TLE calculator [26]. These patients also had multiple leads to be removed (including three or more leads), they were more likely to require venous approach on both sides of the chest, extraction of leads with redundant loops in the heart, extraction of abandoned lead (s) and extraction of lead (s) with long or very long implant duration (Table 2).
Table 2. Patient-, system- and procedure-related risk factors for procedure complexity and major complications.

| No Stenosis 1 | Mild Stenosis 2 | Moderate Stenosis 3 | Severe Stenosis 4 | Total Occlusion 5 | ANOVA Kruskal–Wallis Test (1–5) p | Mann–Whitney U/Chi² Tests (1–2) vs. (4–5) |
|---------------|-----------------|---------------------|-------------------|------------------|-----------------------------------|----------------------------------|
| Mean ± SD n (%) | Mean ± SD n (%) | Mean ± SD n (%) | Mean ± SD n (%) | Mean ± SD n (%) |                                  |                                  |
| **Number of Patients** | | | | | | |
| N = 499 (17.15%) | N = 574 (19.73%) | N = 605 (20.80%) | N = 581 (19.97%) | N = 650 (22.34%) |                                  |                                  |
| **Patient-related risk factors for TLE complexity and complications** | | | | | | |
| Patient age during TLE [years] | 64.48 ± 14.85 | 66.74 ± 13.69 | 66.64 ± 13.79 | 68.57 ± 13.04 | 67.68 ± 14.32 | <0.001 | <0.001 |
| Male gender | 325 (65.13) | 324 (56.44) | 324 (53.55) | 342 (58.86) | 447 (68.77) | <0.001 | 0.019 |
| Etiology: IHD, MI | 278 (55.71) | 329 (57.31) | 353 (58.34) | 361 (62.13) | 355 (54.62) | 0.112 | |
| Etiology: non-ischemic | 221 (44.29) | 245 (42.69) | 252 (41.66) | 220 (37.87) | 295 (45.38) | 0.112 | |
| NYHA class III and IV (%) | 67 (13.42) | 96 (16.72) | 71 (11.73) | 196 (33.74) | 99 (15.23) | 0.523 | |
| LVEF < 40% | 158 (31.66) | 173 (30.14) | 177 (29.26) | 196 (33.74) | 215 (33.08) | 0.847 | |
| Renal failure (any) | 88 (17.64) | 119 (20.73) | 122 (20.17) | 121 (20.83) | 156 (24.00) | 0.211 | |
| Previous sternotomy | 81 (16.23) | 85 (14.81) | 79 (13.06) | 76 (13.08) | 114 (17.54) | 0.210 | |
| Charlson comorbidity index [number of points] | 4.543 ± 3.789 | 4.702 ± 3.380 | 4.688 ± 3.589 | 5.086 ± 3.629 | 4.728 ± 3.550 | 0.038 | 0.016 |
| **CIED system and history of pacing** | | | | | | |
| Device type—pacemaker (any) | 350 (70.14) | 402 (70.04) | 416 (68.76) | 384 (66.09) | 461 (70.92) | 0.193 | |
| Device type—ICD-V, ICD-D | 126 (25.25) | 127 (22.13) | 144 (23.80) | 141 (24.27) | 127 (19.54) | 0.142 | |
| Device type—CRT-D | 21 (4.208) | 44 (7.666) | 44 (7.237) | 56 (9.639) | 62 (9.54) | 0.006 | 0.002 |
| Redundant loop of the lead on X-Rays before TLE | 20 (4.01) | 21 (3.659) | 25 (4.132) | 26 (4.475) | 44 (6.769) | 0.074 | 0.047 |
| Lead with proximal end in CVS before TLE | 9 (1.804) | 10 (1.742) | 7 (1.150) | 11 (1.893) | 14 (2.154) | <0.001 | 0.762 |
| Number of CIED-related procedures before TLE (SD) | 1.728 ± 0.970 | 1.701 ± 0.881 | 1.777 ± 1.068 | 1.806 ± 0.959 | 2.088 ± 1.300 | <0.001 | <0.001 |
| TLE before current TLE | 23 (4.609) | 23 (4.007) | 19 (3.140) | 27 (4.647) | 39 (6.00) | 0.200 | |
| **Risk factors for major complications and procedure complexity** | | | | | | |


Table 2. Cont.

|                         | No Stenosis 1 | Mild Stenosis 2 | Moderate Stenosis 3 | Severe Stenosis 4 | Total Occlusion 5 |
|-------------------------|---------------|-----------------|---------------------|-------------------|------------------|
| **Number of Patients**  | N = 499 (17.15%) | N = 574 (19.73%) | N = 605 (20.80%)   | N = 581 (19.97%)  | N = 650 (22.34%) |
| **Mean ± SD**           | n (%)         | n (%)           | n (%)               | n (%)             | n (%)            |
| Number of extracted leads per patient [n] | 1.597 ± 0.619 | 1.582 ± 0.681 | 1.623 ± 0.666 | 1.683 ± 0.719 | 1.835 ± 0.880 | <0.001 | <0.001 |
| Three or more leads extracted | 30 (6.012) | 46 (8.014) | 55 (9.090) | 73 (12.57) | 113 (17.38) | <0.001 | <0.001 |
| Approach: left          | 483 (96.79) | 551 (95.99) | 578 (95.54) | 548 (94.32) | 599 (92.15) | 0.002 | <0.001 |
| Approach: right         | 7 (1.403)   | 9 (1.568)     | 12 (1.980)         | 17 (2.930)       | 11 (1.690)     | 0.443 |
| Approach: both          | 1 (0.200)   | 3 (0.523)     | 3 (0.496)          | 5 (0.860)        | 14 (2.154)     | 0.003 | 0.009 |
| Approach: femoral       | 1 (0.200)   | 2 (0.348)     | 2 (0.331)          | 0 (0.00)         | 6 (0.923)      | 0.225 |
| Approach: subclavian-femoral | 3 (0.601) | 4 (0.697) | 3 (0.496) | 2 (0.344) | 5 (0.970) | 0.581 |
| Approach: other, combined | 4 (0.802) | 3 (0.523) | 5 (0.826) | 9 (1.549) | 14 (2.150) | 0.061 | 0.017 |
| Approach: Jugular       | 0 (0.00)    | 2 (0.348)     | 0 (0.00)           | 0 (0.00)         | 0 (0.00)       | 0.086 | 0.420 |
| Extraction of leads with redundant loop | 14 (2.806) | 14 (2.439) | 20 (3.306) | 21 (3.614) | 37 (5.690) | 0.026 | 0.011 |
| Extraction of broken lead with proximal end in CS | 11 (2.204) | 11 (1.916) | 7 (1.157) | 10 (1.721) | 16 (2.460) | 0.756 |
| Extraction of abandoned lead(s) (any) | 35 (7.014) | 45 (7.840) | 39 (6.446) | 60 (10.33) | 102 (15.69) | <0.001 | <0.001 |
| Extraction of abandoned lead(s) [n] | 0.074 ± 0.277 | 0.103 ± 0.380 | 0.083 ± 0.340 | 0.138 ± 0.441 | 0.208 ± 0.520 | <0.001 | <0.001 |
| Oldest extracted lead (months) | 96.25 ± 73.71 | 102.4 ± 74.46 | 100.3 ± 76.37 | 94.21 ± 70.22 | 104.86 ± 76.70 | 0.078 |
| Average (per patient) extracted lead dwell time [months] | 92.24 ± 67.47 | 97.85 ± 67.97 | 95.38 ± 70.31 | 88.59 ± 62.52 | 96.49 ± 67.68 | 0.163 |
| Cumulative dwell time of extracted leads [years] | 12.30 ± 12.11 | 13.57 ± 12.43 | 13.55 ± 12.47 | 12.98 ± 11.59 | 15.39 ± 14.13 | 0 < 0.001 | 0.008 |
| SAFeTY TLE calculator of risk for MC [points] | 5.290 ± 4.117 | 5.828 ± 4.130 | 5.995 ± 4.249 | 5.597 ± 4.090 | 6.333 ± 4.560 | 0.002 | <0.024 |
| SAFeTY TLE calculator of risk for MC [%] | 1.470 ± 2.566 | 1.621 ± 2.490 | 1.782 ± 3.430 | 1.608 ± 2.690 | 2.089 ± 3.650 | 0.002 | <0.001 |

CRTD—implantable cardioverter-defibrillator with resynchronization function, CS—coronary sinus, ICD-D—dual chamber implantable cardioverter-defibrillator, ICD-V—single chamber implantable cardioverter-defibrillator, IHD—ischemic heart disease, LVEF—left ventricular ejection fraction, MI—myocardial infarction, NYHA—New York Heart Association functional class, TLE—transvenous lead extraction.
Table 3. TLE complexity.

| No Stenosis | Mild Stenosis | Moderate Stenosis | Severe Stenosis | Total Occlusion | ANOVA Kruskal–Wallis Test (1–5) p | ANOVA Kruskal–Wallis Test (1–2) vs. (4–5) |
|-------------|---------------|-------------------|----------------|----------------|-----------------------------------|------------------------------------------|
| **Number of Patients** | | | | | | |
| N = 499 | N = 574 | N = 605 | N = 581 | N = 650 | | |
| **Mean ± SD** | **Mean ± SD** | **Mean ± SD** | **Mean ± SD** | **Mean ± SD** | | |
| n (%) | n (%) | n (%) | n (%) | n (%) | | |
| **TLE complexity** | | | | | | |
| Procedure duration (skin-to-skin) [minutes] | 57.66 ± 22.54 | 59.18 ± 25.99 | 58.23 ± 20.90 | 59.33 ± 24.09 | 64.16 ± 33.85 | 0.018 | 0.075 |
| Procedure duration (sheath-to-sheath) [minutes] | 12.36 ± 19.06 | 13.26 ± 21.86 | 12.73 ± 17.14 | 13.56 ± 20.36 | 20.32 ± 32.45 | <0.001 | <0.001 |
| Average time of single lead extraction [minutes] | 8.149 ± 11.26 | 8.441 ± 15.05 | 7.755 ± 9.616 | 7.568 ± 8.884 | 10.56 ± 15.30 | <0.001 | 0.126 |
| All leads were extracted | 399 (79.96) | 432 (75.26) | 450 (74.38) | 420 (72.29) | 503 (77.38) | 0.040 | 0.181 |
| Functional lead was left in place for continuous use | 98 (19.64) | 137 (23.87) | 153 (25.29) | 157 (27.02) | 141 (21.69) | 0.030 | 0.207 |
| Non-functional lead was left in place | 1 (0.200) | 3 (0.523) | 1 (0.165) | 1 (0.172) | 6 (0.923) | 0.302 |
| Non-functional superfluous lead was extracted | 35 (7.014) | 45 (7.840) | 39 (6.446) | 60 (10.33) | 102 (15.69) | <0.001 | <0.001 |
| Technical problem during TLE (any) | 85 (17.03) | 110 (19.16) | 116 (19.17) | 109 (18.76) | 162 (24.92) | 0.011 | 0.025 |
| Block in implant vein (subclavian region) | 34 (6.814) | 39 (6.794) | 48 (7.934) | 43 (7.401) | 65 (10.00) | 0.154 |
| Lead-to-lead binding | 28 (5.611) | 33 (5.749) | 39 (6.446) | 42 (7.229) | 64 (9.846) | 0.030 | 0.009 |
| Byrd dilator collapse/torsion/“fracture” | 16 (3.206) | 19 (3.310) | 19 (3.140) | 19 (3.270) | 23 (3.538) | 0.968 |
| Lead fracture during extraction | 22 (4.409) | 22 (3.833) | 31 (5.124) | 29 (4.991) | 51 (7.846) | 0.004 | 0.014 |
| Need to change venous approach | 12 (2.405) | 13 (2.265) | 14 (2.314) | 18 (3.098) | 41 (6.308) | <0.001 | <0.002 |
| Functional lead dislodgement | 6 (1.202) | 8 (1.394) | 6 (0.992) | 5 (0.861) | 5 (0.769) | 0.824 |
Table 3. Cont.

|                          | No Stenosis 1 | Mild Stenosis 2 | Moderate Stenosis 3 | Severe Stenosis 4 | Total Occlusion 5 | ANOVA Kruskal–Wallis Test (1–5) p | ANOVA Kruskal–Wallis Test (1–2) vs. (4–5) |
|--------------------------|---------------|-----------------|---------------------|-------------------|-------------------|-----------------------------------|-------------------------------------------|
| Number of Patients       | N = 499       | N = 574         | N = 605             | N = 581           | N = 650           |                                   |                                           |
| Mean ± SD n (%)          |               |                 |                     |                   |                   |                                   |                                           |
| Loss of lead fragment    | 2 (0.401)     | 8 (1.394)       | 6 (0.992)           | 5 (0.861)         | 5 (0.769)         | 0.800                             |                                           |
| Reel of ICD lead coil    | 2 (0.401)     | 4 (0.697)       | 5 (0.826)           | 1 (0.172)         | 2 (0.308)         | 0.222                             |                                           |
| Number of big technical problems | 1.316 ± 0.637 | 1.325 ± 0.718 | 1.313 ± 0.685       | 1.330 ± 0.620     | 1.500 ± 0.784     | 0.002                             | 0.053                                     |
| One technical problem only | 57 (11.42)    | 65 (11.32)      | 76 (12.56)          | 64 (11.01)        | 81 (12.46)        | 0.925                             |                                           |
| Two technical problems   | 16 (3.206)    | 12 (2.091)      | 12 (1.983)          | 21 (3.614)        | 33 (5.077)        | 0.006                             | 0.029                                     |
| Three or more technical problems | 3 (0.601)   | 6 (1.045)       | 8 (1.322)           | 3 (0.515)         | 14 (2.154)        | <0.001                            | 0.302                                     |
| Other smaller technical problems | 25 (5.010) | 23 (4.007)      | 26 (4.298)          | 22 (3.787)        | 49 (7.538)        | 0.003                             | 0.192                                     |
| Use of additional tools  |               |                 |                     |                   |                   |                                   |                                           |
| Evolution (old and new) or tight rail | 7 (1.403) | 5 (0.871)       | 8 (1.322)           | 7 (1.205)         | 17 (2.615)        | 0.121                             |                                           |
| Metal sheath             | 30 (6.012)    | 36 (6.272)      | 50 (8.264)          | 40 (6.886)        | 63 (9.692)        | 0.064                             | 0.051                                     |
| Lasso catheter/snare     | 14 (2.806)    | 13 (2.265)      | 17 (2.810)          | 17 (2.926)        | 33 (5.077)        | 0.017                             | 0.052                                     |
| Basket catheter          | 7 (1.403)     | 6 (1.045)       | 4 (0.661)           | 2 (0.344)         | 8 (1.231)         | 0.284                             |                                           |

ICD—implantable cardioverter-defibrillator, TLE—transvenous lead extraction.
Table 4. TLE efficacy and complications, and long-term mortality after TLE.

| No Stenosis | Mild Stenosis | Moderate Stenosis | Severe Stenosis | Total Occlusion |
|-------------|---------------|-------------------|-----------------|-----------------|
| N = 499     | N = 574       | N = 605           | N = 581         | N = 650         |

| Mean ± SD | Mean ± SD | Mean ± SD | Mean ± SD | Mean ± SD |
|-----------|-----------|-----------|-----------|-----------|
| n (%)     | n (%)     | n (%)     | n (%)     | n (%)     |

TLE efficacy and complications

| Major complications (any) | 8 (1.603) | 10 (1.742) | 11 (1.818) | 16 (2.754) | 16 (2.461) | 0.599 |
| Hemopericardium           | 5 (1.002) | 6 (1.045)  | 8 (1.322)  | 11 (1.893) | 9 (1.385)  | 0.707 |
| Hemorrhage                | 3 (0.601) | 3 (0.523)  | 1 (0.165)  | 0 (0.00)   | 3 (0.462)  | 0.288 |
| Tricuspid valve damage    | 2 (0.401) | 4 (0.697)  | 2 (0.331)  | 5 (0.861)  | 3 (0.462)  | 0.722 |
| Rescue cardiac surgery    | 4 (0.802) | 3 (0.523)  | 8 (1.322)  | 9 (1.549)  | 10 (1.538) | 0.286 |
| Minor complications (any)| 31 (6.212)| 35 (6.098) | 46 (7.438) | 44 (7.229) | 60 (8.615) | 0.278 |
| Procedure-related death   | 1 (0.200) | 1 (0.174)  | 1 (0.165)  | 2 (0.344)  | 1 (0.154)  | 0.401 |
| Indication-related death  | 1 (0.200) | 0 (0.00)   | 0 (0.00)   | 1 (0.172)  | 0 (0.00)   | 0.485 |
| Partial radiographic success | 13 (2.605)| 18 (3.136) | 22 (3.636) | 18 (3.098) | 34 (5.231) | 0.203 |
| Complete clinical success | 492 (98.60)| 563 (98.08)| 592 (97.85)| 570 (98.11)| 634 (97.54)| 0.806 |
| Complete procedural success| 481 (96.39)| 550 (95.82)| 578 (95.54)| 556 (95.70)| 611 (94.00)| 0.322 |

Organizational model of TLE procedure. TEE monitoring

| Routine TEE monitoring of lead extraction | 261 (52.31) | 266 (46.34) | 263 (43.37) | 263 (45.27) | 233 (35.85) | <0.001 |
| TLE-related TV dysfunction               | 22 (4.409)  | 26 (4.530)  | 29 (4.793)  | 38 (6.540)  | 39 (6.000)  | 0.234  |
Table 4. Cont.

|                           | No Stenosis 1 | Mild Stenosis 2 | Moderate Stenosis 3 | Severe Stenosis 4 | Total Occlusion 5 | ANOVA Kruskal–Wallis Test (1–5) | ANOVA Kruskal–Wallis Test (1–2) vs. (4–5) |
|---------------------------|---------------|-----------------|---------------------|------------------|------------------|--------------------------------|------------------------------------------|
| Number of Patients        | N = 499       | N = 574         | N = 605             | N = 581          | N = 650          |                                |                                           |
| Mean ± SD n (%)           |               |                 |                     |                  |                  |                                |                                           |
| Increase in TR by 2 degrees | 8 (1.603)     | 5 (0.871)       | 13 (2.149)          | 10 (1.721)       | 11 (1.692)       | 0.632                          |                                           |
| Increase in TR by 3 degrees | 2 (0.401)     | 2 (0.348)       | 3 (0.496)           | 0 (0.00)         | 3 (0.462)        | 0.892                          |                                           |
| Increase in TR by 2 degrees and to Grade IV | 2 (0.401) | 4 (0.697) | 3 (0.496) | 5 (0.861) | 3 (0.462) | 0.839                          |                                           |
| Damage to chordae tendineae during TLE | 14 (2.806) | 15 (2.613) | 18 (2.975) | 24 (4.131) | 25 (3.846) | 0.531                          |                                           |
| **Short-, mid- and long-term mortality after TLE** |                     |                 |                     |                  |                  |                                |                                           |
| Survival in 1712 ± 1187 (1–4638) days of follow up | 375 (75.15) | 389 (67.77) | 436 (72.07) | 373 (64.20) | 409 (62.92) | <0.001                         | <0.001                                    |
| Death within 48 h | 3/499 (0.601) | 1/574 (0.174) | 1/605 (0.165) | 3/581 (0.516) | 2/650 (0.308) | 0.794                         |                                           |
| One-month mortality; 2–30 days; n (% of patients with follow-up longer than 2 days) | 4/495 (0.808) | 9/573 (1.571) | 5/604 (0.828) | 4/578 (0.692) | 7/648 (1.080) | 0.464                         |                                           |
| One-year mortality (31–365 days); n (% of patients with follow-up >30 days) | 31/491 (6.314) | 40/558 (7.168) | 43/597 (7.203) | 38/573 (6.632) | 37/640 (5.781) | 0.889                         |                                           |
| Three-year mortality (366–1095 days); n (% of patients with follow-up >365 days) | 24/433 (5.543) | 51/482 (10.581) | 50/514 (9.728) | 74/499 (14.83) | 59/579 (10.19) | <0.001                         | <0.001                                    |
| Death at >3 years (at 1095 days); n (% of patients with follow-up >1095 days) | 62/359 (17.27) | 84/429 (19.58) | 70/447 (18.57) | 89/447 (19.91) | 136/569 (23.90) | <0.001                         | <0.001                                    |

TEE—transesophageal echocardiography; TLE—transvenous lead extraction TR—tricuspid valve regurgitation, TV—tricuspid valve.
Analysis of TLE complexity and degree of LRVSO showed that procedure duration (sheath-to-sheath), extraction of non-functional superfluous leads, occurrence of any technical problem during TLE, lead-to-lead binding, lead fracture during extraction, need to change venous approach, coincidence of three or more technical problems and necessity of using metal sheaths and lasso catheters/snares were associated with the presence of LRVSO (Table 3).

The occurrence of any major complication, urgent rescue cardiac surgery, partial radiographic success (remained tip or <4 cm lead fragment), damage to chordae tendineae, other forms of TLE-related TV dysfunction/damage, complete clinical success and complete procedural success as well as procedure-related death (intra-, post-procedural) did not show any relationship with LRVSO, similar to mortality in the first day, first month and first year after TLE. In contrast, mortality at more than one-year follow-up was significantly higher among patients with severe venous stenosis and complete venous occlusion (Table 4).

Analysis of mortality using the Kaplan–Meier curve confirmed the relationship between LRVSO and long-term survival after TLE (Figure 1).

Figure 1. Probability of survival of patients undergoing TLE depending on the degree of LRVSO.

4. Discussion

Venous obstruction is a well-known complication after implantation of a permanent transvenous pacemaker. The incidence of venous obstruction reaches 30–45% with complete occlusion rates of 12% on average and 1–3% for symptomatic occlusion [1–20]. In the current study, severe venous obstruction was identified in 19.94% (40.77% if moderate occlusion was included) whereas complete occlusion in 22.34% of patients. The higher
incidence rate of total occlusion in the present study may be a result of long implant duration: cumulative dwell time of the extracted leads was 15.31 ± 12,925 years. Closer evaluation of the clinical factors showed that LVRSO was more common in elderly males with a higher Charlson comorbidity index. Several investigators confirm the contribution of various clinical factors to the occurrence of venous complications [4–6], others show no association between LVRSO and the clinical condition of the patient [7,11]. Analysis of the system/procedure-related factors in the present study demonstrated that the number of extracted leads, lead extraction on the left side or both sides of the chest, extraction of the lead with redundant loop in the heart, extraction of abandoned leads, extraction of leads with long implant duration and a higher risk of MC estimated using the SAFeTY TLE calculator [26] were related to the presence of severe venous stenosis or total venous occlusion. LVRSO was also more common in patients with CRT, having leads with their proximal end in the CVS and a higher number of CIED-related procedures before lead extraction. A similar relationship, especially between the number of extracted leads/long implant duration and LVRSO has been shown in previous reports [5,11].

Out of 20 reports, only four described LRVSO diagnosed just before the TLE procedure [2,6,13,14], and only two assessed the influence of LRVSO on the complexity of TLE [2,6]. The last two studies provide contradicting results. Li et al. in a study of 202 patients concluded that the presence of LRVSO made it more difficult to extract the leads, requiring advanced tools and more time [2]. In contrast, Boczar et al. in a group of 133 pts demonstrated that LRVSO did not influence the effectiveness, safety, and the use of additional tools during TLE procedures [6]. In the present study, the indicators of procedural difficulty and complexity such as procedure duration, extraction of superfluous leads, occurrence of any technical problem, lead-to-lead binding, fracture of the extracted lead, need to change venous approach, coincidence of three or more so-called technical problems and need to use metal sheaths or lasso catheters were related to the presence of LRVSO. The occurrence of any major complication was insignificantly higher in groups with LRVSO as compared to groups without significant stenosis: 2.754 and 2.461% vs. 1.603% and 1.742%, respectively. The need to perform urgent rescue cardiac surgery, partial radiographic success and damage to chordae tendinae during TLE were not significantly associated with the degree of LRVSO. The occurrence of TLE-related TV damage, achievement of complete clinical success and complete procedural success as well as procedure-related death (intra-, post-procedural) were unrelated to LRVSO, similar to mortality in the first day, first month and first year after TLE. This study, however demonstrated a link between TLE difficulty/complexity and the degree of LRVSO, which may be a reflection of implant duration and the total number of extracted leads. Thus, the real problem is only with implantation of new lead(s) because of lead dysfunction or necessity of upgrading the CIED system.

The pathophysiology of LRVSO is not well understood. It is likely that lead-related endothelial trauma incites an inflammatory response of the vessel wall with subsequent thrombosis and scarring. Early (days, weeks) LRVSO seems to be a result of thrombosis which can be treated with low-molecular heparin [16–20]. The role of thrombosis in delayed (months) or late (years) LRVSO is less clear. The inflammatory response of the vessel wall probably induces the formation of scar tissue similar to lead adhesion to the vessel and heart structures, observed on the extracted leads and during TEE [33]. The process of natural maturation makes lead-related fibrotic scar harder and harder leading to its mineralization and calcification. It is well-known that scar tissue in the SVC and in the heart makes lead dissection more difficult [34]. However, so far, nobody has considered scar tissue causing LRVSO and scar tissue around the leads detected during TEE/ICS as the same phenomenon. Looking at narrowing or occlusion of implant veins from this viewpoint we can explain the relationship between LRVSO and TLE complexity, difficulty and complications. Lead dissection in scarred veins is more effort-consuming and sometimes requires stronger pulling on the lead to be extracted. It can also explain the mechanism of TV damage during TLE (fortunately rare). It seems to confirm the concept of simultaneous lead traction from
above and below during dissection; it can protect both the SVC wall and the TV [35]. Our results seem to confirm the significance of routine venography before TLE and considering LRVSO as still another risk factor for TLE complexity and major complications.

In the present study, worse long-term survival was demonstrated in patients with a higher degree of LRVSO. The reason for the worse survival rate in this group is not clear and is probably related to other factors as well (possibly a higher Charlson index).

5. Study Limitations

Our study has several limitations worth noting. Routine venography before TLE was performed in all patients except those with contraindications, mainly renal failure. For this reason, an interesting patient subpopulation had been excluded from the study. The database was prospectively integrated, but analysis was performed retrospectively. For the purposes of this study, the population of patients was divided into groups according to maximal venous narrowing without taking into account the site of narrowing/occlusion and the length of venous stenosis/occlusion. Therefore, the present analysis of venograms includes maximal venous narrowing but not the volume of the phenomenon (the number of vessels affected). The classification of patients we used in the study not only enabled comparison of our results with the findings of other investigators, but also maximal venous narrowing was considered a practical marker for predicting the usefulness of veins for implantation of a new lead/catheter.

6. Conclusions

The occurrence of significant venous stenosis/occlusion in patients undergoing TLE is related to some clinical factors (age, male gender, high Charlson comorbidity index) and numerous procedure-related factors, especially long implant duration, extraction of leads with redundant loop in the heart, extraction of abandoned leads, presence of leads with proximal end in the coronary sinus vein and a higher number of CIED-related procedures before lead extraction. LRVSO can be considered as an additional risk factor for TLE complexity. Further research is required to provide evidence for the relationship between scar tissue density encapsulating the leads visible in TEE and the degree of LRVSO. Lead-related venous stenosis/occlusion has no influence on mortality at one-year follow-up, but the presence of severe forms of LRVSO is associated with worse prognosis of patients undergoing TLE at more than one-year follow-up.

Author Contributions: Writing—original draft preparation, M.C.; methodology and statistical study, W.J.; investigation and corresponding author, A.P.; data curation, J.K., D.N., Ł.T. and P.S.; writing-review and editing, A.K. All authors have read and agreed to the published version of the manuscript.

Funding: This research received no external funding.

Institutional Review Board Statement: The study was conducted according to the guidelines of the Declaration of Helsinki, and approved by the of Bioethics Committee at the Regional Medical Chamber in Lublin protocol number 288/2018/KB/VII.

Informed Consent Statement: Informed consent was obtained from all subjects involved in the study.

Data Availability Statement: Readers can access the data supporting the conclusions of the study at www.usuwanieelektrod.pl (accessed on 11 September 2021).

Conflicts of Interest: The authors declare no conflict of interest.

Abbreviations

| Abbreviation | Description                      |
|--------------|----------------------------------|
| AnV          | innominate (brachiocephalic) vein|
| AxV          | axillary vein                    |
| CIED         | cardiac implantable electronic device |
| CRT          | cardiac resynchronization therapy |
| EF           | ejection fraction                |
FU follow-up
ICD implantable cardioverter-defibrillator
IVC inferior vena cava
LR lead-related
LRVSO lead-related venous stenosis/occlusion
LV left ventricle
LVEF left ventricular ejection fraction
NYHA The New York Heart Association (functional class)
Pts patients
PM pacemaker
RA right atrium
RV right ventricle
TEE transesophageal echocardiography
TLE transvenous lead extraction
ScV subclavian vein
SVC superior vena cava
TV tricuspid valve
VSO venous stenosis/occlusion

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