Cardiac implantable electronic device lead extraction using the lead-locking device system: keeping it simple, safe, and inexpensive with mechanical tools and local anesthesia

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ABSTRACT

Objective: We have previously reported our successful approach for percutaneous cardiac implantable electronic device (CIED) lead extraction using inexpensive tools, which we have continued over the years. Herein we report the results of the systematic use of a unique stylet, the lead-locking device (LLD), which securely locks the entire lead lumen, aided with non-powered telescoping sheaths in 54 patients to extract 98 CIED leads.

Methods: This prospective observational clinical study included 38 men and 16 women aged 68.9±13.1 years undergoing lead extraction for device infection (n=46), lead malfunction (n=5), or prior to defibrillator implant (n=3). Leads were in place for 6.7±4.3 years. Infections were more commonly due to Staphylococcus species (n=40). There were 78 pacing (31 ventricular, 37 atrial, 4 VDD, and 6 coronary sinus leads) and 20 defibrillating leads.

Results: Using simple traction (6 leads) and the LLD stylets (92 leads) aided with telescoping sheaths (15 patients), 96 (98%) leads in 52 (96.3%) patients were successfully removed, with all but one leads removed using a subclavian approach; in 1 patient, the right femoral approach was also required. In 2 patients, distal fragments from one ventricular pacing and one defibrillating lead could not be removed. Finally, lead removal was completely (52/54) (96.3%) or partially (2/54) (3.7%) successful in 54 patients for 96 of 98 leads (98%) without major complications.

Conclusion: Percutaneous lead extraction can be successful with mechanical tools using the LLD locking stylet aided with non-powered telescoping sheaths through a simplified, safe, and inexpensive procedure using local anesthesia alone. (Anatol J Cardiol 2017; 18: 289-95)

Keywords: cardiac implantable electronic devices, pacing lead, defibrillating lead, lead extraction, pacemaker infection, lead malfunction

Introduction

The need for lead extraction of cardiac implantable electronic devices (CIEDs), including pacemakers, defibrillators (ICD), and cardiac resynchronization therapy (CRT) devices, is increasing (1) because the number of device implantations and their attendant complications along with the infection burden associated with the implantation are increasing over time (2, 3) and because malfunctioning and recalled leads continue to emerge (4).

Percutaneous extraction of infected or dysfunctional CIED leads has supplanted surgical techniques with great success (5-7). Current methods employ mechanical and/or laser equipment with variable success rates. The locking stylet has remained the principal tool in these techniques, whereas the telescoping (powered or non-powered) mechanical or laser sheaths serve as the most important ancillary tool. Various locking mechanisms have been used, but most of them in addition to requiring tedious and exact sizing of the stylet to match the lead lumen, lock at the tip of the lead, which may not always be readily accessible. Newer types of locking stylets have simplified the sizing process and facilitated and strengthened the locking mechanism. Cognizant of the expense entailed with powered mechanical means, particularly that of the prohibitive cost for our practice of the laser technique, our standard approach over the years has included only mechanical means using a locking stylet, non-powered sheaths, and other ancillary tools, with all procedures performed under local anesthesia. We have previously reported our experience with various types of locking stylets with exclusive distal tip locking (8, 9), and we are herein extending it with use of another type of stylet, the lead-locking device (LLD) (Spectranetics, Colorado Springs, CO, USA), which when locked, occupies the lumen of the lead over its entire length, in a group of 54 consecutive patients who also required some additional tools (mostly non-powered sheaths) for lead extraction, attempting to extract 98 pacemaker, ICD, and CRT leads.
Methods

Study design
This was a prospective observational clinical study.

Patients
Over a period of 10 years, pacing, ICD, or CRT lead extraction using the LLD system was attempted in 54 consecutive patients who were referred to our institutions for percutaneous lead removal; we excluded patients (n=5) for whom different extraction means were employed. Prospective data were recorded for all patients and procedures. The indication for lead removal was device infection (46 patients), pacing (2 patients), or ICD (3 patients) lead dysfunction or the removal was performed prior to an ICD implantation (3 patients) (Table 1). Infections involved both pocket and lead(s) and were mainly due to S. epidermidis (n=34). Skin erosion was present in 16 patients. Positive blood cultures were detected in 9 (19.6%) of 46 patients with infection. Echocardiography revealed small-/moderate-sized vegetations on the right ventricular pacing leads in 4 patients. No patient had evidence of pulmonary embolism. The indication for permanent pacemaker implantation (n=34) was sinus node dysfunction in 9 patients, neurally mediated syndromes in 2 patients, and atrioventricular block in 23 patients. Sustained ventricular tachycardia was the indication for ICD implantation in 14 patients, whereas 6 patients with refractory heart failure had a CRT system implanted.

Type of extracted leads
The specific types of extracted leads can be seen in Table 1. Pacing leads (37 atrial, 35 ventricular, and 6 coronary sinus) had been implanted in 78 patients. Defibrillating leads had been implanted in 20 patients. Leads had been in place for a mean of 6.7±4.3 years (range 0.3–19 years). The fixation mechanism of implanted leads was passive in 92 leads and active (screw-in) in 6 leads. Of the 78 pacing leads, 4 were unipolar ventricular, 21 bipolar ventricular, 6 bipolar CS, 1 unipolar atrial, 36 bipolar atrial, 4 unipolar single-pass VDD, all tined, and 6 bipolar screw-in atrial (n=2) or ventricular (n=4) leads. The ICD defibrillation leads were tined or active fixation double-coil (n=18) or single-coil (n=2) leads.

Locking stylet and telescoping sheaths
The LLD locking stylet comprises an expandable wire mesh woven around a stylet, which when inserted and locked in place, occupies the whole lumen of the lead over the entire length of the stylet body, allowing for traction along the whole lead and not just at its tip (Fig. 1). It was available to us in 2 sizes (LLD#1 and LLD#2). It can accommodate a wide range of leads with inner lumen diameters ranging from 0.017” (0.43 mm) to 0.028” (0.66 mm) for LLD#2. By locking along the entire lead lumen, it provides a stable traction platform. The plastic telescoping sheaths used in this series for the dissection of lead adhesions and to provide counter traction were the ones used in our prior series, the mechanical, non-powered, S and L short Extor sheaths (VascoMed GmbH, Binzen, Germany) with inner diameters of 8.5F and 12.5F, respectively.

Procedure of lead extraction
All procedures were performed after an informed written consent was obtained from each patient. Each case was also discussed with our cardiothoracic surgeon who covered the procedure. For the pacemaker-dependent patients, a temporary pacing wire was inserted usually from the internal jugular vein or occasionally from the contralateral subclavian vein. Then, with use of sterile and aseptic technique and local anesthesia,
For patients with an infected system, intravenous antibiotic therapy was continued for 4–6 weeks as appropriate; a new device system was implanted, usually on the contralateral side, after 10–14 days had elapsed and when the device pocket was clean of any signs of infection and laboratory testing indicated that inflammatory or infection indices had returned to normal. For patients with non-infected systems, a new system was implanted either during the same session or the next day. After hospital discharge, all patients were followed up at the pacemaker clinic and by their own referring physicians.

Statistics
The data are expressed as mean±standard deviation (range) and/or percentages. Comparisons were made using the Student’s unpaired t-test for quantitative data and chi-square analysis and z-statistic for qualitative and proportional data. Correlation between variables was calculated using the Pearson correlation coefficient. Data were analyzed with SPSS 23 (Armonk, NY). A p value of <0.05 was considered significant.

Results
Procedural outcome
In preparation for the extraction procedure, a temporary pacemaker electrode catheter was inserted in 8 (14.8%) patients who were pacemaker dependent. Lead extraction was attempted in 54 patients for a total of 98 leads (78 pacing and 20 ICD defibrillating leads). Complete removal of all leads was successful in 52 (96.3%) patients for 96 (98%) leads; partial lead removal with retention of a lead fragment was obtained in 2 patients. Both patients with retained ventricular lead fragments were patients with device infections: 1 with a single-chamber ventricular pacemaker and a lead dwell time of 15 years and the other with an atrioventricular ICD device implanted 13.4 years earlier. The former patient did well conservatively, responding to antibiotic therapy, whereas the other patient preferred elective surgery over a transfemoral approach for the removal of the retained ICD lead fragment. The LLD#1 stylet was used in 2 patients and the LLD #2 stylet in 52 patients. Successful removal was attained using a right (n=28) or left (n=24) or right plus left (n=1) subclavian or right subclavian plus femoral (n=1) approach. Lead extraction was accomplished by simple traction for 4 atrial, 1 ventricular, and 1 coronary sinus lead (only test stylet inserted), sole use of the locking stylet for 60 (47.4%) leads in 39 (58%) patients, locking stylet aided by non-powered sheaths for 27 leads, and a femoral approach for 1 ventricular lead. The procedures lasted for 1.2±0.9 h (range 20 min to 5 h), with a fluoroscopy time of 9.0±6.7 min.

Use of a non-powered telescopic sheath
In addition to the locking stylet, telescoping sheaths to aid lead extraction were required for 27 (27.6%) leads in 15 patients (27.8%) (Fig. 2a, b; Table 2). The transfemoral approach was needed in 1 patient. In 1 patient with a dual chamber pacemaker and 2 bipolar leads, the stylet was used successfully to remove the...
atrial lead using a right subclavian approach, and a bioptome and a snare were used to capture and remove the ventricular lead via a transfemoral approach after attempts from above had failed (Fig. 2c). Coronary sinus leads were successfully removed in 6 (6.1%) patients (Fig. 2d); telescoping sheaths were used in 2 patients.

Transfemoral approach

As already indicated, in 1 patient with a dual-chamber pacemaker, a right subclavian approach was successful in removing the atrial lead with use of the LLD stylet aided by a sheath, but the ventricular lead could not be extracted using the approach from above and thus a transfemoral approach was used for this purpose. The lead was severed proximally, a bioptome was used to capture and release the proximal end of the lead, and then a snare (Noose catheter, VascoMed GmbH, Binzen, Germany) was used to successfully extract this lead (Fig. 2c). Coronary sinus leads were successfully removed in 6 (6.1%) patients (Fig. 2d); telescoping sheaths were used in 2 patients.

Complications

No major complications occurred in this series of patients during the procedures of percutaneous lead extraction. A moderate-sized hematoma developed in the pocket of 2 patients who had extensive local debridement. Vagotonia occurred in 2 patients; in one of them, ventricular ectopy with frequent extrasystoles and runs of non-sustained ventricular tachycardia were also observed during lead traction. The latter patient was the one who was submitted to a femoral approach to remove the ventricular lead and this was complicated by femoral vein trauma, which was repaired with suturing.

Correlations

The duration (r=48, p=0.0002) and fluoroscopy time (r=0.54, p=0.0001) of the procedure correlated with the lead dwell time; both parameters were longer in males than in females (p<0.05). Also, patients requiring a telescoping sheath had longer lead dwell time (105.3±46.2 months vs. 72.3±51.4 months; p=0.035). Finally, the use of a mechanical sheath incurred longer procedure (1.7±1.2 h vs. 0.9±0.6 h; p=0.004) and fluoroscopy times (13.1±8.8 min vs. 7.6±5.1 min; p=0.006).

Discussion

The results of the present study indicate that percutaneous lead extraction can be successful with mechanical tools using the LLD stylet owing to its unique locking mechanism aided by non-powered telescoping sheaths through a simplified, safe, and inexpensive procedure using local anesthesia alone. Thus, we were able to completely extract 96 of 98 (98%) CIED leads, with a mean implant time of 6.7 years. To the best of our knowledge, this is the second series of patients reported in the
literature with the systematic use of the LLD system. In a prior report of the initial multicenter series of 57 patients using the LLD device for extraction of 99 leads combined with a variety of sheaths, 97/99 (98%) leads were successfully removed with no major complications (10).

Implantation of CIEDs has markedly increased over the past several years due to expanded indications (2, 11-13). Device systems include pacemakers, ICDs, and CRT systems (13-15). Unfortunately, complications associated with CIEDs have also been increasing (3, 4, 16). Particularly, device infections appear to be rising out of proportion to increasing implant rates (17). Furthermore, device recipients who develop device infection have increased device-dependent, long-term mortality even after successful treatment of infection (18). The only way to deal with this grisly reality is to take extra measures to prevent or minimize CIED infections by performing a lege artis procedure (19). Thus, in parallel to the increasing complications, there is a growing need for lead extraction to manage such complications (1). Device infection was the main indication for lead extraction in 47 of 54 (87%) patients in this series referred to our centers for the extraction procedure, wherein 9 (19%) of these had developed lead endocarditis.

Extraction of pacemaker and ICD leads has been performed via percutaneous techniques, which have virtually supplanted the surgical approach (1). Lead extraction is quite challenging, particularly for leads implanted for longer periods of time (5, 6). This is because endocardial leads develop encapsulation by fibrous tissue not only at the distal tip, which is in contact with the endomyocardium, but also along the endovenous or endocardial course of the lead(s), where fibrous adhesions impede the extraction (20). Hence, special tools are required to pave the way and facilitate lead extraction. When dealing with infection, the presence and size of vegetation may pose further risk to percutaneous methods (21).

Among the techniques employed, the most widely used lead extraction system is the Cook retrieval system (Cook Vascular, Inc., Vandergrift, PA) in addition to a variety of other tools and systems (5). Success is higher for physicians with greater experience, shorter implant duration, active fixation, and atrial leads. A femoral approach is required in a small percentage of patients. However, there were problems with the availability of these old styles; thus, we subsequently resorted to the LLD type, which admittedly has an easier and more effective locking mechanism that involves the entire lead body and length (Fig. 1). Furthermore, we adopted a more frequent use of beveled sheaths to facilitate the extraction process, particularly for leads of a longer implant duration. These measures appeared to obviate the need to resort to a transfemoral approach, which was needed in only 1 patient in the current series, and also avoided the need to use an array of ancillary tools required in 27% in our previous experience (9). Most importantly, the use of the LLD system was safe with no major complications observed during the procedures. Based on our prior experience, persistence of using only traction via the other types of locking styles was probably responsible for the occurrence of cardiac tamponade in a paraplegic patient with severe left ventricular dysfunction and infective endocarditis (not included in this series), who subsequently required cardiac surgery but finally succumbed to a low cardiac output state.

With regard to coronary sinus leads, no difficulties were encountered in 6 patients undergoing lead removal in our series. Other series have also confirmed the feasibility of extracting such leads without any additional problems that were initially anticipated, except for a certain type of active fixation leads (31).

Study limitations

This study was not a randomized controlled trial. It was rather our experience in our lead extraction program based on a relatively small series of patients. However, the data were pro-
spectively collected for all patients. The study included all consecutive patients in whom the LLD system was utilized but not all consecutive patients undergoing lead extraction because there were few other patients during the study period depending on tool availability, in whom other extraction means were employed. A potential drawback of our extraction approach may be associated with the unavailability of a powered mechanical sheath system, which most likely led to a relatively long procedural time (4–5 h) in 3 patients in the current series. Finally, although we found the LLD stylet easier to use and rather more effective than its predecessors, we cannot comment on the relative merits of the other commonly used locking stylet (Liberator by Cook Medical) because we have only recently started using it.

Conclusion

The LLD lead extraction system with a uniform whole lumen/entire lead length applicable locking stylet aided by mechanical sheaths appears to be simple to operate, safe, and quite successful in chronically implanted lead extraction and can be applied without the need of general anesthesia through a low-cost approach. Further studies in larger patient populations are needed to confirm these results and randomized comparative, and cost-effectiveness studies are necessary for the currently available lead removal systems.

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References


