Demonstration of proof-of-concept of StrokeShield system for complete closure and occlusion of the left atrial appendage for non-valvular atrial fibrillation therapy

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Abstract

In the US, the most significant morbidity and mortality associated with non-valvular atrial fibrillation (NVAF) is embolic stroke, with 90% of thrombus originating from the left atrial appendage (LAA). Anticoagulation is the preferred treatment for the prevention of stroke in NVAF patients, but clinical studies have demonstrated high levels of non-compliance and increased risk of bleeding or ineligibility for anticoagulation therapy, especially in the elderly population where the incidence of NVAF is highest. Alternatively, stroke may be prevented using clinically approved surgical and catheter-based devices to exclude or occlude the LAA, but these devices continue to be plagued by peri-device leaks and thrombus formation because of residual volume. To overcome these limitations, Cor Habere (Louisville, KY) and the University of Louisville are developing a LAA closure device (StrokeShield) that completely occludes and collapses the LAA to minimize the risk of stroke. The StrokeShield device is a collapsible occluder (nitinol reinforced membrane) that completely covers the LAA orifice with an expandable conical coil anchor that attaches to the myocardium. The device is designed for catheter-based delivery and expands to completely occlude the LAA orifice and collapse the LAA. The primary advantages of the StrokeShield system are a completely sealed LAA (no peri-device flow or residual space) and smooth endothelialized connection to the left atrial wall with minimal risk of cardiac bleeding and tamponade. We tested proof-of-concept of a prototype StrokeShield device in acute (n = 2) and chronic 60-day (n = 2) healthy canine models. Acute results demonstrated that the conical coil securely attached to the myocardium (5N pull-out force) and the Nitinol umbrella fully deployed and covered the LAA ostium. Results from the chronic implants demonstrated long-term feasibility of device placement with no procedural or device-related intra- or post-operative complications, secure placement and correct positioning of the device with no device migration. The device successfully occluded the LAA ostium and collapsed the LAA with no interference with the mitral valve, circumflex coronary artery, or pulmonary veins. Necropsy demonstrated no gross signs of thrombus or end-organ damage and the device was encapsulated...
Foundation to fund the subaward (PI: GG) to develop the prototype StrokeShield device and conduct proof-of-concept testing, which are presented in this manuscript. The funders (Wallace Coulter Foundation) had no role in the study design, data collection and analysis, decision to publish, and/or preparation of the manuscript.

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MS, SK, JJ are co-founders and LT is an engineer with Cor Habere (Louisville KY) who are collectively continuing to develop the technology (StrokeShield system) presented in this manuscript. Note: all of the research and development presented in this manuscript was completed before Cor Habere was incorporated. Cor Habere was also awarded a NIH SBIR phase I grant (R43HL142337-01, PIs: JJ and MS and co-Is: GM, SK, GG) to continue development work of StrokeShield system, but none of research and development funded by the SBIR phase I grant is completed before Cor Habere was incorporated. Cor Habere was also awarded a NIH SBIR phase I translation-partnership-award-in-biomedical-engineering-tp/ to fund the subaward (PI: Giridharan). The Coulter Foundation (non-profit) provided a 5-yr Clinical Translational Research award to the University of Louisville to be distributed as subawards to University of Louisville faculty (clinicians, engineers, scientists) to develop proof-of-concept medical device(s) and/or technology(s) awarded through a well-defined internal and external peer-review process. The project presented in this manuscript was supported with funding as one of these selected subawards (PI: Giridharan). The Funder (Coulter Foundation) provided funding support for materials and supplies only (no salary support for any of the authors), but did NOT have any role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript. The specific roles of the authors are articulated in the ‘authors contributions’ section.

Competing interests: Financial - Two US patents awarded related to the research presented in this manuscript, and a third University of Louisville (UofL) invention disclosure was filed February 1, 2021, but is not presented in this manuscript. Slaughter MS, Giridharan GA, Sobieski MA, Soucy KG, and Koenig SC. Atrial appendage closure device and related methods (US Patent 10,531,878 awarded January 14, 2020) Slaughter MS, Giridharan GA, Sobieski MA, Soucy KG, and Koenig SC. Atrial appendage closure device and related methods (US Patent 10,898,202 awarded January 26, 2021) Slaughter MS, Monreal G, Koenig SC, Jimenez J, Tompkins L. Left Atrial Appendage (LAA) device, delivery tool, and related methods (UofL Invention Disclosure no. 21038, filed in the LAA. Histology demonstrated mature neointima covering the device with expected foreign body inflammatory response. These early positive results will help to guide the iterative design process for the continued development of the StrokeShield system.

Introduction

The incidence of nonvalvular atrial fibrillation (NVAF) is increasing worldwide and represents a major health care burden [1, 2]. Despite advances in medical care, prognosis for NVAF remains poor due to the risk of embolic stroke from flow stagnation, especially in the left atrial appendage (LAA). In the US, there are 2.6 to 6.1 million patients with NVAF, which is expected to increase to 12 million by 2050 [3, 4]. The treatment of NVAF is estimated to cost between $16 and $26 billion annually [3, 5], with AF-related hospitalizations accounting for over half of these costs (52%) [5]. The most significant morbidity and mortality associated with NVAF is embolic stroke, with 90% of thrombus originating in the LAA [6, 7]. NVAF is associated with a five times greater risk of ischemic stroke, and NVAF is the main contributing factor for up to 25% of strokes in patients over the age of 80 years [8].

Current treatment options to reduce the occurrence of strokes related to NVAF include medical and surgical therapies. The use of anticoagulation, although effective in preventing stroke, has been associated with bleeding as well as challenges with patient noncompliance (e.g. taking their medications) [9, 10]. Alternatively, a number of mechanical devices have been developed that are designed to prevent thrombus by occluding or excluding the LAA. Current AHA/ACC Guidelines recommend the use of percutaneous LAA occlusion devices for patients who are poor candidates for anticoagulant therapy due to propensity for bleeding or poor drug tolerance and/or adherence [11]. While the procedure is conceptually simple, development of devices that provide effective anatomic fit and placement (device-ostium shape match) and reliably achieve complete occlusion (no leak) is challenging due in part to the variability of LAA shape and ostium dimensions [12, 13].

To overcome these challenges, the StrokeShield system is being developed, which is comprised of a LAA closure device that fully occludes the LAA orifice and collapses the LAA chamber, independent of orifice geometry and without obstructing the pulmonary veins or mitral valve, and a transseptal catheter-based delivery tool. StrokeShield is designed to completely collapse and seal the LAA, which may help lead to further reductions in the incidence of thromboembolic stroke and adverse events and may potentially enable earlier weaning from anticoagulation. In this article, the basic concept, prototype design, and proof-of-concept testing of the StrokeShield system is presented.

Materials and methods

Device design

The StrokeShield system (US patents 10,898,202, 10,531,878) features of an expanding circular umbrella closure device to occlude the LAA ostium and a coil anchor to secure and collapse the LAA wall for closure of the LAA with complete seal (tissue integration) and no residual chamber space (eliminate LAA volume/prevent peri-device leak), Fig 1. The umbrella is constructed from a single Nickel-Titanium (Nitinol) tube with an expanding lattice design to provide structural integrity and is covered with Dacron to facilitate tissue in-growth and encapsulation. The umbrella is designed to completely cover the LAA ostium and will be
available in multiple sizes (21, 28, 33mm) with oversize fit (20% larger than LAA orifice) to meet the expected range of patient LAA geometries and sizes (16-36mm) [14, 15]. The super-elastic Nitinol design enables the LAA closure device to be pre-collapsed in the forward axial direction of the delivery catheter, which will enable all of the implant devices (independent of size) to be deployed using the same delivery tool via a steerable 12Fr sheath. A conical coil cut into the opposing end of the Nitinol tube is designed to secure the LAA closure device to the LAA wall. The collapsible LAA closure device (umbrella) and coil anchor are pre-loaded on a delivery tool. The anchor is secured to the LAA free wall by a clinician applying rotational force using the hand-held control knob to embed the coil anchor. The 2.5-turn coil provides 35mm$^2$ of anchoring surface area designed to have greater pull-out force than suture in cardiac tissue thereby reducing the risk of device migration or myocardial tear. The coil configuration and geometry are designed to compress the LAA wall by creating an outward tissue dimple on the external surface of the LAA wall due to radial myocardial compression. The anchoring coil design is based upon the successful history of other proven implantable medical device technologies that also use screw-in coil mechanisms (i.e. pacing leads [16], apical closure devices [17]) with an established track record of minimal leaks, superior strength, and the ability to be easily retrievable. Only a single contact point is required to secure the Dacron-covered umbrella to the LAA wall, occlude the ostium, and collapse the LAA wall. This approach is designed to reduce the risk of bleeding or tamponade, but will require extensive pre-clinical testing to demonstrate efficacy and safety.

This study was carried out in accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The protocol was approved by the Institutional Animal Care and Use Committee at the AAALAC-accredited University of Louisville (OLAW/PHS Assurance No. A3586-01).

**Acute in-vivo testing.** Two acute non-survival experiments were performed in a healthy canine model to demonstrate intraoperative proof-of-concept of the StrokeShield LAA closure device.

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**Fig 1. StrokeShield device.** The two primary components of the StrokeShield device (22 mm width, 6 mm height) include conical anchor which screws into the LAA myocardium to secure the device, and a Dacron-covered Nitinol umbrella membrane that unfolds following deployment to occlude and collapse the LAA. The upper (A1-A4) and lower (B1 = B4) photos show device with bare Nitinol wire and Dacron-covered umbrella frame, respectively, for isometric (A1,B1), front (A2,B2), back (A3,B3), and side (A4,B4) views.

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device. Two healthy male dogs (34kg, mongrel hounds, Oak Hill Genetics, Ewing IL) underwent ketamine-diazepam sedation followed by endotracheal intubation for mechanical ventilation under general isoflurane anesthesia. Introducers were placed in the carotid artery and jugular vein for drug/fluid administration, blood draws, and cardiac access. A right thoracotomy was performed, and a purse string suture was placed in the LA dome. Heparin was administered and a needle and wire were inserted into the LA dome. The entry point was serially dilated to accommodate the deployment sheath. The device was placed and deployed into the LAA ostium with secure positioning documented by transesophageal echocardiography (TEE) and fluoroscopy. The device remained in situ for a minimum of four hours intraoperatively, after which the animals were euthanized under general anesthesia (IV Beauthanasia D-special, 1mL/10lbs) and necropsies were performed. At necropsy, a device pull-out test was performed.

**Chronic in-vivo testing.** Two 60-day chronic survival experiments were performed in a healthy canine model to evaluate long-term proof-of-concept of the StrokeShield LAA closure device. Two healthy male dogs (27kg, mongrel hounds, Oak Hill Genetics, Ewing IL) underwent LAA device placement as described above. Following device placement, the animals’ thoracotomy incisions were closed, a thoracic catheter was placed for the first post-operative day, the neck lines were removed, post-operative analgesia (fentanyl patch) was initiated, and the animals were extubated and recovered from anesthesia. The animals were maintained post-operatively as needed on aspirin (81mg oral to maintain platelet count within 200–500 K/μL and/or in response to oozing or bleeding at the incision site) and/or Coumadin (2mg oral to maintain INR (international normalized ratio) target range of 2–3). Sixty days later, the animals returned to the fluoroscopy suite and were anesthetized as described above for terminal TEE and fluoroscopic imaging. Following image acquisition, the animals were euthanized under general isoflurane anesthesia (IV Beauthanasia D-special, 1mL/10lbs) and necropsies were performed. Tissue samples were collected for Hematoxylin and Eosin (H&E) and elastic trichrome staining.

**Results and discussion**

**Results**

**Acute in-vivo testing.** Intraoperative placement of the StrokeShield device via the LA dome approach was demonstrated. Acute results demonstrated that the conical coil securely attached to the myocardium and the Nitinol umbrella fully deployed and covered the LAA ostium. At necropsy, up to 5N (510 gram) of pull-out force was applied to the device without tearing LAA tissue or extracting the device demonstrating secure attachment of the coil in the LAA myocardium.

**Chronic in-vivo testing.** Long-term feasibility of device placement with no procedural or device-related intra- or post-operative complications was demonstrated. The animals demonstrated normal cardiac and neurological exams across the 60-day study. Both TEE and fluoroscopy (Fig 2) demonstrated secure placement and correct positioning of the device with no device migration or peri-device flow. The device successfully occluded the LAA ostium and collapsed the LAA with no interference with the mitral valve, circumflex coronary artery, or pulmonary veins (Fig 3). There was no evidence of device/coil perforation or gross LAA cardiac tissue injury or pericardial effusion at implant, during the chronic 60d test period, or at terminal necropsy. Necropsy demonstrated no gross signs of thrombus or end-organ damage. The device was encapsulated in the LAA. Histology using Hematoxylin and Eosin stain (H&E) and Elastic Trichrome stain was performed. Findings demonstrated healed and mature neointima covering the Dacron and Nitinol struts of the closure device. Mature fibrovascular
connective tissue filled the closure device cavity and occlusion point at the contact between closure device disc and atrial wall with normal healing. Mature neointima covered the closure device and mature fibrocellular neointima lined the Dacron cover with expected minimal to mild foreign body response (Fig 4) [18].

Discussion

Between 2004–10, the estimated prevalence of atrial fibrillation (AF) in the US was 4.3 million with nearly 700,000 undiagnosed cases (13%), and its occurrence increases with age [19]. In their review of the epidemiology of AF, Kronej et al. report that as survival with chronic diseases in the elderly population has improved the incidence and prevalence of AF continues to increase, prompting them to suggest AF is becoming a “global epidemic” [20]. Currently, the commercial landscape in the NVAF field of mechanical device therapies, including Watchman™, Amplatzer™, Occlutech™, WaveCrest™, LAmbré™, and Ultrasel™ devices, indicating a robust interest by clinicians and their patients in the continued development and propagation of LAA mechanical device technologies. The Watchman is the only FDA approved LAA closure device with PROTECT [21] and PREVAIL [22] randomized clinical trials demonstrating non-inferiority to oral anticoagulant (warfarin) for ischemic stroke protection but not overall efficacy. Recently, a meta-analysis comparing Watchman and Amplatzer devices for stroke protection resulted in similar low complication and event rates findings with comparable efficacy and safety results [23]. There are also many other emerging LAA mechanical device technologies in various stages of pre-clinical development [24–28].
LAA occlusion devices are designed to block and/or fill the LAA ostium, which if not completely occluded, can result in blood leakage and stagnation near the exposed surrounding edges of the LAA orifice increasing the potential risk for thrombogenesis (and stroke). LAA occlusion devices are designed to block and/or fill the LAA ostium, which if not completely occluded, can result in blood leakage and stagnation near the exposed surrounding edges of the LAA orifice increasing the potential risk for thrombogenesis (and stroke).

Fig 3. StrokeShield device in situ. The StrokeShield device in the LAA at necropsy (day 60) in the chronic canine model demonstrating tissue encapsulation and freedom from interference with the mitral valve and pulmonary veins.

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Fig 4. Histology performed on 60-day chronic canine myocardium and StrokeShield device. At left, Elastic Trichrome (ET) stain demonstrating healed and mature neointima covering the Dacron and nitinol struts (dark black shapes) of the closure device (black dotted line), fully mature fibrovascular connective tissue filling the closure device cavity (red asterisk), and occlusion point at the contact between closure device disc and atrial wall with normal healing (red arrowhead). The black box is further detailed in the upper right image (ET stain). Here, the dotted line indicates healed and mature neointima covering the closure device with irregular edges and microtears indicative of handling artifact. The black double arrow is mature fibrocellular neointima lining the Dacron cover. At lower right, Hematoxylin and eosin stain (H&E) (lower right) showing healed and mature neointima (dotted line) covering the closure device with irregular edges and microtears indicative of handling artifact. The black double arrow shows mature fibrocellular neointima lining the Dacron cover and the single arrow shows Dacron with expected minimal to mild foreign body inflammatory response [18].

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devices with membrane-covered frames may only partially fill the LAA chamber (leaving residual volume) with potential risk for developing a large thrombus within the LAA cavity following occlusion. The LAA tissue, geometric shape, and size of the LAA ostium varies from patient-to-patient making accurate sizing and fixation of the implant challenging as well as risk of the device perforating and/or causing tissue injury. The incidence of pericardial effusion requiring surgery, need for pericardiocentesis, and device embolization have been reported to be 0.4–1.6%, 1.5–2.9%, and 0.4–0.7%, respectively, in clinical trials [29]. Despite the clinical limitations and technological challenges of LAA mechanical devices, there is strong clinical interest and justification for their continued development and advances field [30].

The StrokeShield system combines the advantages of LAA exclusion (surgical) and occlusion (catheter-based delivery) devices into a single LAA closure procedure by completely collapsing the LAA between the ostium (umbrella blocks opening) and free wall (conical coil) with a secure anchoring mechanism. The unicity of the StrokeShield system is its hybrid concept (LAA collapse and occlusion), advancement of technology (patent-protected anchor and umbrella design and methods), and potential clinical benefits, including eliminating need for anti-coagulation post-implant). The StrokeShield device deploys a conical coil that anchors the device to the LAA wall. The axial length of the conical coil provides up to 35 mm² cross-sectional area within the LA wall to achieve greater than 5N pull-out load while providing a strong and secure single-point attachment to the LAA free wall to reduce the risk of device migration. The axial length of the coil is designed to control depth of penetration into LAA wall (number of coils, conical shape) and provide LAA tissue compression to reduce the risk of pericardial effusion and cardiac tamponade. The StrokeShield collapsible occluder device is projected to be fabricated in different patient-specific sizes (21, 28, 33 mm) and implanted in sizes ~20% larger than the LAA orifice and curved toward the LA wall to completely cover the LAA orifice regardless of orifice geometry without obstructing the pulmonary veins or mitral valve. Next design iteration(s) of the collapsible occluder device will be developed for delivery using a steerable, multi-stage catheter delivery tool (12Fr) through femoral vein access. The catheter delivery tool will be designed for advancement through the venous vasculature into the right atrium (RA), curved using a steerable component to allow for transeptal access into the LA, and then used to anchor and deploy the collapsible occluder to completely cover and occlude the LAA ostium and collapse the LAA to eliminate chamber volume and flow.

To demonstrate proof-of-concept, prototype StrokeShield devices were tested in healthy acute and 60-day canine models. In the acute experiments, surgical approach with access for delivery and deployment in the LAA was achieved demonstrating proper anatomic fit for closure and exclusion of the LAA ostium without tearing tissue or causing cardiac tamponade. In the chronic experiments, preliminary evaluation of device efficacy and biocompatibility was achieved as evidenced by elimination of LAA residual volume and peri-device flow, and no device migration, perforation, or inflammation. Several challenges associated with prototype devices were identified and engineering solutions for the next series of design iterations are being carefully considered. Specifically, the methodology for pre-loading device into the catheter sheath, proper sizing for a wide range in LAA size and variability in LAA shape, promotion of healthy device encapsulation and endothelialization, reducing the risk for coil perforation, tissue damage, and/or inflammation, and development of a multi-stage delivery tool that enables steering, device retrieval, re-positioning and re-deployment, and prevents over-torquing during device implant are warranted.

**Conclusion**

Proof-of-concept of the StrokeShield system was demonstrated in acute and chronic animal experiments as evidenced by successful implant procedure and proper anatomic fit with no
device migration or perforation, and complete closure and occlusion of the LAA ostium with tissue encapsulation and absence of residual volume and peri-device leak. The continued development of the StrokeShield system may lead to an alternative catheter-based approach for percutaneous delivery of a novel LAA device (Dacron covered Nitinol umbrella) that is secured to the LAA wall via a single anchoring point (Nitinol conical coil) to mitigate the risk of stroke originating from the LAA in patients with NVAF.

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