Thoracic Artificial Lung Design

by

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ABSTRACT

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by

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Chair: Keith E. Cook

Currently there is no sufficient bridge to lung transplant for patients with end-stage lung disease. Thoracic artificial lungs (TAL) are being developed for this purpose. TALs are attached to the pulmonary circulation, and thus their blood flow is provided by the right ventricle (RV). Current TALs possess blood flow impedances greater than the natural lungs, resulting in low cardiac output (CO) when implanted in series with the natural lung or in parallel under exercise conditions. In series attachment is desired so that the natural lung will still filter blood and perform its non-respiratory functions. However, in parallel attachment may allow for high flow applications. The goal of this research was to design a device with minimal impedance which does not cause a significant decrease in CO when attached in series, or in parallel with cases of high device flows, such as exercise. This was done both through the examination of geometry changes to the TAL housing and through the use of a compliant housing material.
A new compliant TAL (cTAL) was designed, prototyped and tested both *in vitro* and *in vivo*. First, computational fluid dynamics (CFD) and fluid-structure interaction (FSI) modeling were used to investigate inlet and outlet expansion and contraction angles, $\theta$, of 15°, 45°, and 90° in both hard-shell TALs and cTALs. The 45° model was chosen for the cTAL prototype and tested *in vitro* and *in vivo* in the acute setting, attached both in parallel and in series with the native lungs.

The combination of a gradual entrance and exit to the device, as well as a compliant housing resulted in a device impedance of 0.5 mmHg/(L/min), much lower than the native lungs and all other existing TAL designs. The fiber bundle of the cTAL provided excellent gas transfer, with a rated flow well above 7 L/min. The cTAL developed with this research is capable of lower flow PA-PA attachment, with up to 50% of CO to the cTAL. PA-LA attachment of the cTAL will allow for excellent exercise tolerance and unloading of the RV in patients with pulmonary hypertension.
Chapter 1
Introduction

Motivations and Objectives

The only long term solution for chronic lung disease is lung transplantation; however, organ donation is limited and cannot supply the demand. Currently there is no sufficient bridge to lung transplant for patients with end-stage lung disease. Available devices and support methods are unable to provide complete respiratory support for these patients without causing blood damage or further deterioration of the disease state, often resulting in multiple organ failure. Thoracic artificial lungs (TAL) are being developed without blood pumps for these patients in order to minimize blood damage, coagulation, and inflammation. TALs are attached to the pulmonary circulation, and thus their blood flow is driven by the right ventricle (RV).

Several groups, including our lab, have developed a TAL for long-term respiratory support which is either placed in parallel or in series with the natural lungs. In vivo testing has revealed weaknesses in all existing TAL device designs, including high device impedance which can cause performance problems in both attachment configurations. Current TALs possess blood flow impedances greater than the natural lungs, resulting in low cardiac output (CO) when implanted in series with the natural lung. However, in series attachment is desired so that the natural lung will still filter
blood and perform its non-respiratory functions. Recent studies in our lab with the MC3 Biolung which simulate rest and exercise conditions have shown advantages in placing the device in parallel. CO was maintained with high flows to the TAL in rest and ambulatory conditions, albeit with significantly elevated right ventricular pressure. The goal of this research was to design a device with minimal impedance which does not cause a significant decrease in CO when attached in series, or in parallel with cases of high device flows, such as exercise. This was done both through the examination of geometry changes to the TAL housing and through the use of a compliant housing material.

A new compliant TAL (cTAL) was designed, prototyped and tested both in vitro and in vivo. First, computational fluid dynamics (CFD) and fluid-structure interaction (FSI) modeling were used to examine different housing geometries in both hard-shell TALs and cTALs. Second, in vitro hemodynamic testing was conducted with the TAL and cTAL models. Gas transfer performance of the cTAL was also assessed in vitro. Finally, the optimal cTAL prototype was tested in vivo in the acute setting, attached both in parallel and in series with the native lungs.

**Lung Disease and Transplantation**

Lung disease is the third leading cause of death in the United States, responsible for 1 in 6 deaths. These death rates are currently increasing, and more than 35 million Americans have chronic lung diseases [1]. There are four categories of chronic lung disease which lead to end-stage lung disease: obstructive lung disease (asthma, chronic obstructive pulmonary disease (COPD)), restrictive lung disease (interstitial lung
disease), infectious lung disease (cystic fibrosis), and pulmonary vascular disease (pulmonary arterial hypertension). There are many therapies used to treat these diseases, but the only permanent cure is lung transplantation. COPD, idiopathic pulmonary fibrosis (IPF), cystic fibrosis, emphysema, and idiopathic pulmonary arterial hypertension (IPAH) are the most common diseases that lead to lung transplantation [2,3]. However, there is a shortage of donor lungs available for transplant in these patients. In the U.S. in 2009, there were 1,660 lung transplantations but 2,280 new patients registered on the waiting list [3].

**Respiratory Support Options**

When a patient’s lungs are unable to adequately maintain blood O₂ or CO₂ requirements, respiratory support is required. There are several different respiratory support techniques and devices, some of which are still under investigation. One of the most widely used is mechanical ventilation.

Mechanical ventilation can be used for breathing support in both acute and chronic diseases (Figure 1.1).
The mechanical ventilator applies pressure to a patient’s airway through an endotracheal or tracheostomy tube, causing gas to flow into the lungs until the ventilator breath is terminated. Airway pressure drops to a set baseline expiratory pressure, causing elastic recoil of the chest wall and passive exhalation. Acute respiratory distress syndrome (ARDS) is respiratory failure due to lung inflammation and increased vascular permeability caused by direct or indirect lung injury. Current treatment of ARDS involves mechanical ventilation with positive end expiratory pressure (PEEP). The goal of mechanical ventilation is to support a patient’s breathing while allowing his/her lungs to recover. However, ventilation can also cause direct damage to the lungs. Complications from ARDS include infections, pneumothorax, and lung scarring, all

Figure 1.1: Illustration of a patient on a mechanical ventilator.

which can be caused by ventilation [4]. Deaths from ARDS usually results from multiple system organ failure (MSOF). Research suggests mechanical ventilation triggers the release of inflammatory mediators into the lungs and circulation, a mechanism termed biotrauma, possibly causing MSOF [5]. Advances in ventilator strategies have reduced ventilator-induced lung injury and mortality in ARDS patients [6]. Mechanical ventilation could also be used for a short period to support respiratory function in chronic lung disease patients. However, in 2011 the International Society for Heart and Lung Transplantation reported ventilation prior to transplant as a significant risk factor for both 1-year and 5-year mortality in adult lung transplant recipients [2]. Thus, mechanical ventilation is a contraindication for lung transplantation.

There are several alternatives to mechanical ventilation which provide either partial or total respiratory support. Extracorporeal membrane oxygenation (ECMO, also known as extracorporeal life support) is used to treat cardiac or respiratory failure providing respiratory support (V-V ECMO) or respiratory and cardiac support (V-A ECMO) for days or weeks. ECMO is prolonged extracorporeal (outside the body) cardiopulmonary support through vascular cannulation. In V-V ECMO, blood is taken from the venous system, routed through the circuit where it is oxygenated, and then returned to the venous system (jugular vein in and out, or jugular vein to right femoral vein). For V-A ECMO, blood is taken from the venous system and returned to the arterial system (carotid artery and vein, femoral artery and vein, or aorta and right atrium). The ECMO circuit is large, consisting of: a blood pump, a servoregulator, an oxygenator, a heat exchanger, cannulas, tubing, and continuous heparin infusion (Figure 1.2).
ECMO is used more frequently in neonates in whom survival rates are higher than in adults [7]. The ECMO circuit has a large priming volume and utilizes a pump which causes continuous mechanical stress resulting in blood cell damage [8]. Like mechanical ventilation, ECMO has been considered to be a contraindication to lung transplant with a low 1-year survival rate of 45% [9,10]. Several retrospective studies report the clinical use of ECMO to successfully bridge patients to lung transplant; however, these studies have varying post-transplant survival rates that are dependent on initial patient selection [11,12]. Initial examination of the use of ECMO in awake, non-intubated patients (no mechanical ventilation support) shows improved 1-year survival rates and a more
promising bridging technique [13]. Advances in pump and oxygenator technology and using ECMO as an alternative to mechanical ventilation have improved outcomes and shown the possibility of using ECMO as a bridge to lung transplantation. However, it is still an invasive, complex technique requiring a team of specialists and blood transfusions. Because of its complexity and number of components, there are many avenues for complications. Complication rates are reported by the ELSO registry, broken down by type and mode of support [14]. Some of the most common are: oxygenator failure (8-17%), blood clots in the system components (9-15%), cannula problems (5-15%), and bleeding at the surgical site (6-33%) or cannula site (7-21%).

Arterio-venous carbon dioxide removal (AVCO₂R) is another respiratory support technique which uses a simple arterio-venous (AV) shunt (i.e. femoral artery to vein) with a gas exchanger (Figure 1.3).
CO\textsubscript{2} is transferred across the membrane of the gas exchanger, while nearly all O\textsubscript{2} diffuses across the native lungs. Blood flow is driven by the patient’s arterial blood pressure, eliminating the need for a pump. Device flows up to 25\% of CO are achieved while up to 96\% of CO\textsubscript{2} produced is removed; allowing reduction of ventilator support to 16\% of baseline minute ventilation [15]. This technique has been used clinically to treat patients with ARDS and pneumonia [16-18]. In these studies, hypercapnia was reversed, allowing for less aggressive ventilator settings. The disadvantage to this therapy is low O\textsubscript{2} transfer due to oxygenated blood entering the device and, to a lesser extent, the low blood flows.

Figure 1.3: Arterio-venous CO\textsubscript{2} removal circuit. Blood flows from a cannula inserted in the femoral artery, through a membrane oxygenator where gas exchange occurs, and then back to the patient through a cannula inserted in the femoral vein.

Therefore, AVCO₂R benefits patients with severe hypercapnia and ARDS, but not patients with hypoxemia [8]. More recently, AVCO₂R has been utilized for a bridge to lung transplant [11, 19]; however, oxygenation was limited due to blood flow restrictions and thus could only be used in patients with severe hypercapnia and only moderate hypoxia. Also, only patients with adequate mean arterial pressure (>70 mmHg) and sufficient cardiac output could undergo this procedure. Thus, this technique is not adequate as a bridge to lung transplant.

The only clinical device for AVCO₂R is the Novalung iLA (Figure 1.4).

![Novalung iLA](image)

Figure 1.4: Novalung iLA. Spillner J et al. *Frontiers in Bioscience* 16: 2342-2351, 2011

This device has also recently been used as a bridge to transplant, connected between the pulmonary artery and left atrium. This has been initially used in patients with end-stage pulmonary hypertension [20-23]. Patients were supported anywhere from 8-62 days until transplantation. During that time, Schmid et al reported that pulmonary artery pressure decreased down to 45-60 mmHg; however, this still constitutes rather severe pulmonary hypertension. The device was exchanged, on average, every 15 days due to fibrin deposition which decreased gas exchange. In all these studies, device flow ranged from
1.8-3.0 L/min during support. This attachment mode allows the patient to be mobile; however, with increasing physical activity CO increases. Schmid et al reported that as their patient resumed physical activity, increased CO led to more flow through native lung since flow to device couldn’t be increased. Thus, a larger fraction of unsaturated blood arrived at the left atrium. The resistance of the Novalung is approximately 5-6 mmHg/(L/min) at blood flows of 2-2.5 L/min [17,18,24]. The Novalung was not intended for PA-LA use when constructed. A device with lower resistance that allows for higher flows would provide total respiratory support and allow patients to be ambulatory and even exercise while awaiting lung transplant.

Intravascular oxygenators, small devices implanted in the vena cava, have been studied for more than 20 years and offer advantages over ECMO which include a small blood contact surface, decreased operation time, and a priming volume of zero. However, clinical trials with one such device, the IVOX, showed inadequate gas exchange, providing O₂ transfer of only 20-30% of whole body requirements [25,26]. This device consists of a bundle of hollow fibers which are joined together at their ends. The IVOX is inserted, through the femoral vein, in a compact state and the fibers are then unfurled in the vena cava (Figure 1.5(a)). O₂ flows through the center lumen of the device and then through the fibers (Figure 1.5(b)). Gas exchange takes place between the lumen of fibers and the circulating blood.
Recent advances in device design have been made to increase blood flow through the device with the incorporation of a small axial pump, thus increasing gas exchange (Figure 1.6).

Figure 1.5: (a) Picture of the IVOX in its unfurled state and (b) cross-sectional schematic of the IVOX, showing the gas flow path through the device. Kallis et al. *Eur J of Cardiothorac Surg* 7(4): 206-210, 1993.

Figure 1.6: Highly integrated intravascular membrane oxygenator (HIMOX) utilizing a series of disc-shaped fiber bundles along a core and an axial blood pump. Cattaneo *et al. ASAIO J* 52:180-5, 2006.
Gas exchange was also increased by decreasing fiber porosity, however this led to a higher pressure drop resulting in bundle deformation and a blood shunt around the device [27]. Another method of increasing blood velocities and enhancing gas exchange is the utilization of a pulsating balloon in the center of a bundle of fibers (Figure 1.7).

Figure 1.7: Schematic of the intravenous respiratory assist catheter, consisting of a pulsating balloon surrounding by a bundle of hollow fibers. Federspiel et al. ASAIO J 42:M435-42, 1996.

The fiber bundle is constrained around an un-inflated balloon, making the device smaller than the vessel lumen size and allowing blood to shunt past the device to reduce flow resistance. By rapidly pulsating the balloon, blood flows through the fiber bundle and passes by the fibers as relatively higher velocities. This intravenous respiratory assist catheter, also known as the Hattler catheter, has been shown to enhance gas exchange over the IVOX, providing O₂ transfer rates of 175-305 mL/min/m², depending on balloon pulsation rate [28]. Investigators predict that a 0.5m² device could provide 50% of total O₂ requirements; however, the devices tested in this study had a surface area of about
0.23m². Since intravascular oxygenators are small, bundle surface area must be maximized to maximize gas transfer. However, this results in higher blood flow resistance, reducing venous return to the heart. Thus, despite the above design improvements, these devices cannot provide total respiratory support and are not a sufficient bridge to lung transplant.

Artificial lung systems which utilize a pump or rotating fiber bundle to drive blood flow have been developed by several groups [29-31]. These rotating fiber bundle devices use centrifugal force and high shear stress to mix the blood and enhance gas exchange (Figure 1.8).

![Figure 1.8: Schematic of the paracorporeal assist lung (PRAL) which utilizes a rotating fiber bundle to enhance gas exchange. Svitek et al. ASAIO J 51(6): 773-80, 2005.](image-url)
The disadvantage of these devices is the high shear rates which cause platelet activation and hemolysis. Previous research demonstrated that platelet activation is a function of shear stress magnitude and duration of application and that shear-induced platelet activation occurs only above a threshold of 12-15 dyne/cm$^2$ [32-34]. Svitek et al estimated duration of support for their device to be 7-10 days. They reported high levels of hemolysis at the higher rotating rates and calculated the maximum shear stress of their device due to rotation to be 80 dynes/cm$^2$. This is well above the threshold for shear-induced platelet activation. Recent testing with a new artificial pump-lung (APL) device has shown little to no hemolysis during 30-day in vivo testing [35]. In this device, blood is drawn from the patient (i.e. right atrium or femoral vein) into the pump chamber. Blood is propelled through a diffuser section, driven by a magnetically levitated rotating centrifugal pump impeller, and then enters the fiber bundle in the radial direction. Gas exchange occurs as blood passes through the bundle, until it finally exits the device and returns to the patient (i.e. pulmonary artery or femoral artery) (Figure 1.9).
This device has the advantage of being compact, with a small priming volume; however it is only designed to provide blood flows of 3.5 L/min. Additional testing may be required to examine device performance in ambulatory or exercising patients and patients with pulmonary hypertension.

**Thoracic Artificial Lung Development**

The devices and techniques described above are either not suitable as a bridge to lung transplant or too early in their development. Such a device must be durable, simple to operate, provide total respiratory support, minimize blood damage and activation, and have low resistance. This device must also allow patients to be awake and ambulatory while awaiting lung transplant.
Thoracic artificial lungs (TAL) are being developed to meet these requirements. TALs do not use a pump, but rather are attached to the pulmonary circulation with the right ventricle (RV) driving blood flow. Elimination of the need for a blood pump greatly reduces blood damage, coagulation, and inflammation. The patient’s own cardiac output (CO) controls the device, making it simple to operate. Depending upon patient requirements, the TAL can be attached in parallel or in series with the natural lungs (Figure 1.10).

During in parallel attachment blood flow is routed from the pulmonary artery (PA), through the TAL and then returned to the left atrium (LA). This configuration reduces impedance of the TAL and natural lung system; however, less blood flows through the TAL. Therefore, some blood flow travels to the systemic circulation without being filtered by the natural lungs, allowing embolic material to reach the major organs.

During in series attachment, blood is routed from the proximal PA through the TAL and then back to the distal PA where the blood then flows through the native lungs and finally back to the LA. This configuration eliminates the risk of systemic emboli by allowing the
lung to perform its filtration and non-respiratory functions. In addition, the entire cardiac output can flow through the TAL, allowing for complete gas exchange. Studies on TAL attachment mode have shown parallel TAL attachment to decrease pulmonary system impedance and increase CO, while in series attachment increases pulmonary system impedance and decreases CO [36].

The MC3 Biolung (MC³, Ann Arbor, MI) has been tested in our lab for up to 30 days in sheep [37,38]. In this device, venous blood flows in through the center of the device, radially across the cylindrical fiber bundle, and exits through the duel outlets. Gas flows through the fibers while blood flows over the fibers, with gas exchange occurring through the fiber walls (Figure 1.11).

![Blood flow path through the MC3 Biolung.](image)

The Biolung was attached in the PA-LA configuration without any coating to enhance biocompatibility. In the 30-day study, the average device use was 9.5 days with 51% of CO flowing to the TAL. Average device resistance over 30 days was 2.6 mmHg/(L/min) at 2 L/min of device flow. The Biolung is now moving toward clinical trial. Recent studies with the MC3 Biolung which simulate rest and exercise conditions have also
shown advantages in placing the device in parallel [39]. CO was maintained with high
flows to the TAL in rest and ambulatory conditions. At exercise conditions, CO
decreased with increasing flow to the TAL. Further decreases in device impedance are
necessary for increased exercise tolerance.

Despite promising Biolung results, in series attachment would still be desired if it
did not induce RV dysfunction. However, current TALs possess blood flow impedances
greater than the natural lungs, resulting in low CO when implanted in series with the
natural lung [40]. Increased pulmonary system impedance in the in series configuration is
caused by the additive impedance of the TAL and natural lung. This induces low CO via
increased RV oxygen demand, decreased RV oxygen supply, decreased contractile
function, and pulmonic valve dysfunction [41-43]. Patients with respiratory failure,
specifically those suffering from PH, may not be able to tolerate any additional
impedance from the TAL. In PH the PA pressure is high, causing increased work for the
right heart and often results in RV failure. Therefore, in series attachment of the TAL will
further increase pulmonary system impedance and the right heart workload, possibly
hastening RV failure.

Impedance is the opposition to pulsatile blood flow and is determined from a
relationship between the time-varying components of pressure and flow. Fourier
transforms are used to convert pressure, $P(t)$, and flow, $Q(t)$, waveforms in the time
domain into a series of sinusoids at their respective harmonics frequencies:

$$P(t) = P_0 + \sum_{n=1}^{N} P_n \sin(\omega_n t + \phi_n)$$

$$Q(t) = Q_0 + \sum_{n=1}^{N} Q_n \sin(\omega_n t + \phi_n)$$
where $P_0$ and $Q_0$ are the average pressure and flow magnitudes. The sinusoid at each harmonic, $n$, has its own magnitude ($P_n$ or $Q_n$), frequency ($\omega$), and phase shift ($\phi$). Harmonic frequencies are equal to integer multiples of the heart rate. Impedance is defined as the ratio of the pressure magnitude to flow magnitude at each harmonic:

$$Z_n = \frac{P_n}{Q_n}$$

The zeroth and first harmonic impedance moduli ($Z_0$ and $Z_1$), are most physically relevant as the opposition to steady flow and flow at a frequency equal to the heart rate, respectively. Since CO is affected mostly by $Z_0$ and $Z_1$, these are our focus. Impedance is a function of TAL resistance, compliance, and inertance, with impedance increasing with increasing resistance and inertance and decreasing with increasing compliance. An inlet compliance chamber (Figure 1.12) has been shown to reduce $Z_1$ by nearly 80% and to a lesser extent, $Z_0$ in both \textit{in vitro} and \textit{in vivo} studies [40,44,45].

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{Figure_1.12.jpg}
\caption{Biolung with a polyurethane compliance chamber attached to its inlet. McGillicuddy \textit{et al.} \textit{ASAIO J} 51:789-794, 2005.}
\end{figure}
McGillicuddy et al minimized Z₀ and Z₁ \textit{in vitro} at compliances greater than 1 and 5 ml/mmHg, respectively. By reducing device impedance, the work of the RV is also reduced by dampening the pulsatile flow and creating a more steady flow throughout the cardiac cycle. Despite reducing device impedance, overall system impedance (Z₀=5.75 mmHg/(L/min) and Z₁=2.41 mmHg/(L/min)) was still greater than that of the natural lungs (baseline Z₀=2.87 mmHg/(L/min) and Z₁=0.96 mmHg/(L/min)) during in series attachment, causing a decrease in CO. Sato \textit{et al} suggest that most of the RV dysfunction is due to an increase in the zeroth harmonic impedance, Z₀, due to its effects on average system pressures, and a smaller decrease in CO is caused by the first harmonic impedance modulus, Z₁, due to its effect on systolic pressures. Kuo \textit{et al} confirmed that Z₀ is the key determinant of RV output in their study examining the effect of pulmonary system impedance on CO [46]. For every 1 mmHg/(L/min) increase in Z₀ there is an approximate 3.65% decrease in CO. Likewise, for every 1 mmHg/(L/min) increase in Z₁ there is a 0.08Z₀% decrease in CO. Akay \textit{et al} performed a similar study, examining the relationship between pulmonary system impedance on CO in sheep with pulmonary hypertension [47]. Results again indicated that Z₀ had a significant effect on percent change in CO, %ΔCO, and found that: %ΔCO = -7.45*ΔZ₀. Therefore, the foremost goal of TAL design should be to reduce Z₀, with a secondary requirement to limit increases in Z₁. Sato \textit{et al} concluded that the resistance of the compliance chamber, device or anastomoses must be reduced to improve CO [40]. The compliance chamber design has a limited volume and a large expansion and contraction which significantly adds to the system resistance, and thus a compliant device is proposed.
Compliant TAL

An entire compliant device, having a housing made out of a flexible material, can accommodate a larger change in volume with less stretching. This was examined through a compliant TAL (cTAL) design which utilized a compliant Biospan (segmented polyurethane) housing (Figure 1.13).

Figure 1.13: Original cTAL prototype, consisting of a compliant Biospan housing and fiber bundle. Cook et al. *ASAIO J* 51: 404-411, 2005.

Initial studies with this device showed a significant dampening of the flow pulse; however the device had excessive inlet and outlet resistances [48]. The inlet and outlet regions had abrupt expansions and contractions, leading to minor losses due to flow separation. Recirculations are created in these regions, decreasing the efficiency of the system and leading to higher device impedance. Uneven blood flow can negatively affect biocompatibility, activating blood clotting factors and causing clots to form in low flow areas. We propose the incorporation of more gradual expansions and contractions in the cTAL to reduce losses, improve flow and reduce resistance in the inlet and outlet housing.
regions. This will be accomplished by decreasing the inlet and outlet expansion angle, $\theta$, of the inlet and outlet housing manifolds (Figure 1.14).

Figure 1.14: Inlet and outlet expansion angle, $\theta$, of the cTAL inlet and outlet housing manifolds.

Decreasing $\theta$ results in the elongation of the entrance and exit to the cTAL, creating more gradual expansions and contractions (Figure 1.15).

Figure 1.15: cTAL models with $\theta=15, 45,$ and $90^\circ$

**Summary of the Study**

The goal of this work was to design a low impedance cTAL which prevents large decreases in CO when used in series with the natural lungs and has excellent exercise tolerance when used in a PA-LA configuration. This was accomplished using CFD and in
vitro models to determine ideal housing geometry. Hard-shell TALs were considered first to determine the effects of housing geometry alone on impedance. FSI analysis was then used to assess a compliant housing’s ability to further reduce impedance in different cTAL models. The ideal cTAL was prototyped and underwent hemodynamic and gas transfer testing in vitro. Finally, cTALs were tested in vivo, both in parallel and in series, with different percentages of the CO diverted to the cTAL.
References


Chapter 2

Thoracic Artificial Lung Housing Design

Introduction

Current TALs and cTALs possess blood flow impedances greater than the natural lungs, resulting in abnormal pulmonary hemodynamics when implanted in series with the natural lung or in parallel with high flows to the TAL [1,2]. Previous studies have shown the majority of cTAL impedance was due to excessive inlet/outlet impedances [3]. It was theorized that the abrupt expansions and contractions at the inlet and outlet regions cause flow recirculation that, in turn, leads to higher device impedance. If this is the case, more gradual expansions and contractions could improve flow patterns and reduce device impedance. Device design also influences flow uniformity within the cTAL. It is essential to have even flow throughout the fiber bundle and housing in order to have efficient gas transfer and eliminate any low flow areas where blood clots can form. Thus, device design is essential to optimize both the impedance and biocompatibility of the cTAL.

The effect of housing geometry on cTAL impedance will be explored in Chapter 3 using fluid-structure interaction (FSI) modeling. However, the influence of housing geometry alone will first be considered in simplified, rigid-housing TAL designs. In this chapter, the inlet and outlet expansion angle, $\theta$, of the TAL was adjusted, and flow patterns and impedance were determined at a variety of blood flow rates and heart rates.
First, computational fluid dynamics (CFD) was used to examine the effects of different housing designs on device flow patterns and impedance. Flow patterns are important in predicting gas transfer performance and biocompatibility and cannot be easily visualized in vitro. Second, physical models were created and tested in vitro to verify computer results.

Methods

Computational Fluid Dynamics Studies

The effects of $\theta$ on TAL blood flow patterns and impedance were evaluated using designs with $\theta=15^\circ$, $45^\circ$ and $90^\circ$ (Figure 2.1).

![Figure 2.1: TAL model showing blood flow path and inlet and outlet expansion angle, $\theta$](image)

SolidWorks (Dassault Systèmes SolidWorks Corp, Concord, MA) computer-aided design (CAD) software was used to create each housing model, which was then imported into the ADINA (ADINA R&D Inc., Watertown, MA) computational fluid dynamics (CFD) software program (Figure 2.2).
In these models, blood flows through the inlet, expands into the inlet manifold, flows through the fiber bundle at the center of the device, then travels through the outlet manifold and exits through the device outlet. The inlet/outlet diameter, height and width of each TAL model are 0.016 m, 0.103 m and 0.102 m, respectively. The fiber bundle length, path length (the distance blood must flow through the bundle), and frontal area (the cross-sectional area perpendicular to blood flow through the bundle) are 0.140 m, 0.038 m, and 0.016 m², respectively, for each model. The 15°, 45°, and 90° models have a total length of 0.350 m, 0.229 m, and 0.191 m. Since the inlet/outlet tube diameter and the housing body geometry is fixed, the length of the expansion/contraction section increases as θ decreases (see Figure 2.1).

**CFD: Problem Formulation**

Program assumptions included a transient solution with incompressible flow. Turbulent flow was assumed to occur in the inlet and outlet sections. The average Reynolds numbers in the inlet and outlet sections at flow rates of 2, 4, and 6 L/min are 920, 1800, and 2760, respectively. Peak Reynolds numbers at these same flows are 3000, 6100, and 9200, respectively. The material properties of the rigid housing and fiber
bundle, along with the fluid characteristics of blood were defined in the program. Blood viscosity and density were set to 0.003 Pa s (3.0 cP) and 1040 kg/m³. The fiber bundle region was modeled as porous media with fluid density, porosity, permeability, and fluid viscosity defined as 1040 kg/m³, 0.75, $2.81 \times 10^{-9}$ m², and 0.003 Pa s (3.0 cP), respectively. Permeability, $k$, was calculated from preliminary cTAL data [3] using Darcy’s Law:

$$ k = \frac{Q \mu L}{A \Delta P} $$  \hspace{1cm} (1)

where $Q$ is blood flow rate, $\mu$ is viscosity, $L$ is the fiber bundle path length, $A$ is the fiber bundle frontal area, and $\Delta P$ is the pressure drop across the fiber bundle. The hollow fiber bundle was modeled using a porous media theory to determine flow characteristics within the bundle [4-7].

The fluid motion outside the fiber bundle is governed by the Navier-Stokes equations with constant density and viscosity, together with the continuity equation as follows:

$$ \rho_f \frac{\partial \mathbf{V}}{\partial t} + \rho_f (\mathbf{V} \cdot \nabla \mathbf{V}) = -\nabla P + \mu_f \nabla^2 \mathbf{V} $$ \hspace{1cm} (2)

$$ \nabla \cdot \mathbf{V} = 0 $$ \hspace{1cm} (3)

where $\rho_f$ is the blood density, $\mu_f$ is the blood viscosity, $P$ is the pressure and $\mathbf{V}$ is the velocity vector.

**CFD: Boundary Conditions**

A symmetry condition was utilized in ADINA, allowing the model to be cut in half, thus reducing computational time. This was done by applying a zero velocity fixity
perpendicular to the symmetry face. A no-slip wall boundary condition was also applied to the inlet and outlet housings, while a slip wall boundary condition was applied to the bundle walls. The momentum equation used in the porous media model does not contain any second order derivatives, thus viscous effects are negligible and a slip wall condition is used. The initial pressure condition was set to 10 mmHg. A time dependent velocity was applied to the inlet face and a constant pressure of 10 mmHg was applied to the outlet face to simulate left atrial pressure.

Pulsatile blood flow through the device was simulated in ADINA at average flow rates ($Q$) of 2, 4, and 6 L/min and heart rates ($HR$) of 80 and 100 beats/min (Figure 2.3).

Figure 2.3: TAL inlet flow waveform for CFD and in vitro studies at 4 L/min, 100 beats/min, and pulsatilities of 2 and 3.75.
This inlet flow waveform, $Q_i$, is a simulated RV output with a pulsatility of 3.75. Pulsatility is defined as (maximum flow rate – minimum flow rate)/average flow rate. The inlet waveform was created with the equation [8]:

$$Q_i = \begin{cases} 
AV_s e^{-t} \sin \left( \frac{\pi t}{t_{es}} \right) & 0 < t < t_{es} + 0.02 \\
Q_s(t_{es} + 0.02) \left[ 1 - \frac{t - t_{es} - 0.02}{0.02} \right] & t_{es} + 0.02 < t < t_{es} + 0.04 \\
0 & t_{es} + 0.04 < t < t_{ed}
\end{cases} \quad (4)$$

where $V_s$ is the stroke volume defined as:

$$V_s = \frac{Q}{HR} \quad (5)$$

$t_{es}$ is the end-systolic time defined as:

$$t_{es} = 0.43 - \left(1.8 \times 10^{-3}\right)HR + \left(4.44 \times 10^{-6}\right)HR^2 \quad (6)$$

$t_{ed}$ is the end-diastolic time defined as:

$$t_{ed} = \frac{60}{HR} \quad (7)$$

and $\lambda$ is the flow pulse scaling factor used to scale flows to match the CO. The duration of regurgitation is assumed to be 0.04s. In reality, device inflow is less pulsatile since the TAL is attached to the pulmonary artery (PA) which dampens the pulse. Thus, this waveform represents the most extreme case the device would encounter. A more typical inlet waveform with a lower pulsatility of 2 was also investigated (Figure 2.3) [3]. This waveform is created by sending the previously discussed RV output waveform through a windkessel model with $R = 1.25 \text{ mmHg}/(L/\text{min})$ and $C = 1.78 \text{ mL/mmHg}$. 

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**CFD: Numerical Scheme**

A finite element formulation based on the Galerkin method was utilized to solve the governing equations. In this method, the governing equations (equations 2 and 3) are converted into discretized equations which can be solved numerically. In order to achieve converged and accurate results, a value of $10^{-5}$ was set for the relative error. The mesh size and time step of the model were varied until a converged solution was reached. The convergence criterion was set at $\leq 1.5\%$ difference in impedance between models of different time step or mesh size. A variable grid size system was employed in the present study to capture the rapid changes in the velocity and pressure variables occurring in specific regions. A finer mesh was implemented at the inlet and outlet faces in order to accurately determine pressure and flow data. Also, a finer mesh along the inlet and outlet cylindrical region was required for convergence. In the end, the mesh size was 3 mm for the bodies, 0.25 mm for the inlet and outlet sections, and 1.0 mm for the inlet and outlet faces. A time step of 0.005 s was used.

**CFD: Data Analysis**

The simulation was run for three consecutive heart beats to achieve a converged periodic solution. Inlet and outlet pressure and overall flow rate data from the final beat were obtained from the program for presentation and analysis. To determine the impedance moduli, Fourier transforms were used to convert the inlet pressure and flow in the time domain to a series of sinusoids at their respective harmonic frequencies:

$$P(t) = P_0 + \sum_{n=1}^{N} P_n \sin(\omega_n t + \phi_n)$$  \hspace{1cm} (8)
\[ Q(t) = Q_0 + \sum_{n=1}^{N} Q_n \sin(\omega_n t + \phi_n) \quad (9) \]

where \( P_0 \) and \( Q_0 \) are the average pressure and flow magnitudes, \( P_n \) and \( Q_n \) are the pressure and flow magnitudes at the \( n \)th harmonic, \( \omega_n \) is the frequency at the \( n \)th harmonic, \( t \) is time, and \( \phi_n \) is the phase shift at the \( n \)th harmonic. Impedance moduli at the \( n \)th harmonic, \( Z_n \), are the ratio of the amplitudes of pressure and flow at the same harmonic:

\[ Z_n = \frac{(P_i - P_o)_n}{Q_n} \quad (10) \]

where \( P_i \) and \( P_o \) are the inlet and outlet device pressures. Values of \( Z_0 \) and \( Z_1 \) were calculated and used to assess TAL impedance. Pressure and flow data were also obtained from the top and bottom fiber bundle faces to determine bundle resistance:

\[ R = \frac{P_{TB} - P_{BB}}{Q} \quad (11) \]

where \( P_{TB} \) is the average pressure across the top fiber bundle face, \( P_{BB} \) is the average pressure across the bottom fiber bundle face, and \( Q \) is average flow through the bundle.

In order to separately investigate the viscous pressure drop, the average pressure drop due to an expansion or contraction, \( \Delta P_{ec} \), was calculated:

\[ \Delta P_{ec} = \frac{1}{(t_2 - t_1)} \int_{t_1}^{t_2} \rho Q(t)^2 \left[ \frac{1}{A_1} - \frac{1}{A_2} \right]^2 dt \quad (12) \]

where \( \rho \) is the density of blood, \( Q(t) \) is the time-dependent blood flow through the device at \( Q=4 \) L/min and \( HR=100 \) beats/min, \( A_1 \) is the cross-sectional area of the inlet/outlet, and \( A_2 \) is the cross-sectional area of the fiber bundle. Since our goal was to understand the relative contributions to total device resistance of each section, and flow didn’t seem to
change this, this calculation was done at one representative flow condition for both the 45° and 90° models. Equation 12 was numerically integrated using the trapezoidal rule. The viscous resistance was then calculated for the inlet \(R_{v, in}\) and outlet \(R_{v, out}\) sections:

\[
R_{v, in} = \frac{\Delta P + \Delta P_{sc}}{Q}, \quad (13)
\]

\[
R_{v, out} = \frac{\Delta P - \Delta P_{sc}}{Q}, \quad (14)
\]

where \(\Delta P\) is the pressure drop for the inlet, bundle, or outlet section, and \(Q=4\) L/min. No adjustment is necessary for the fiber bundle resistance, as it does not include an expansion or contraction. The percentage of total device resistance was then calculated for the inlet, bundle and outlet sections.

Velocity band plots were created in ADINA to investigate flow uniformity in the housing and fiber bundle of each model. Plots were created on a cross section along the y-axis (top down view) for several sections in the housing and fiber bundle of the device. In these plots, color is used to indicate velocity magnitude where red represents the highest velocity in the plot range and blue represents a velocity of 0 m/s. These plots were created at peak systole and end-diastole in the cardiac cycle for two different velocity ranges. Velocity vector plots were also created to investigate areas of recirculation in each housing model. In these plots, color and the size of the vectors are used to indicate velocity magnitude where red represents the maximum velocity in the plot range and blue represents a velocity of 0 m/s.
In Vitro Testing

Experimental tests were run on two TAL rigid housing models to validate the computer results. The models tested had $\theta = 45^\circ$ and $90^\circ$. To create the fiber bundle, woven mats of polypropylene fibers with a fiber diameter of 200 $\mu$m were wound into compact bundles with porosity, path length, and frontal area of 0.75, 0.038 m, and 0.016 m$^2$, respectively. The bundle was potted with WC7-53 (BJB Enterprises Inc., Tustin, CA), a two part polyurethane, into an acrylic box. Three sets of housings for each $\theta$ were machined from PVC blocks and then clamped above and below a fiber bundle. The blocks were carved out in the center, so that they are the negative of the housing shown in Figure 2.1. In vitro testing was conducted with a circuit consisting of a pulsatile pump (Harvard Apparatus, Holliston, MA), TAL model, and reservoir. The entire circuit was primed with a 3.0 cP glycerol solution (Figure 2.4).

![Figure 2.4: In vitro test circuit consisting of a pulsatile pump, TAL model, and reservoir filled with 3.0 cP glycerol solution. Six pressure lines were connected to the model at the inlet (P1), outlet (P2), top of the bundle (P3 and P5), and bottom of the bundle (P4 and P6).](image1)

Pressure transducers (see Figure 2.4) were connected to six luers on the model in order to monitor device pressure at the inlet (P1), outlet (P2), top of the bundle (P3 and P5), and bottom of the bundle (P4 and P6). A blood flow probe was placed before the
device inlet to monitor flow rate. Using the pulsatile pump, testing was conducted for $HR= 80$ and $100$ beats/min and $Q= 2, 4, \text{ and } 6$ L/min. Average pulsatilities for these flows were 3.33. A representative inlet flow rate waveform at a $HR=100$ beats/min, $Q=4$ L/min, and pulsatility = 3.33 is shown in Figure 2.3. In addition, the $45^\circ$ model was tested at a pulsatility of 1.7 with $HR=100$ beats/min and $Q=4$ L/min to examine the effects of pulsatility on device impedance. To reduce the pulsatility, a compliance chamber [2] with $C = 0.5$ was placed in the circuit before the device inlet. The BIOPAC MP150 data acquisition system (Biopac Systems, Inc., Goleta, CA) was used to acquire pressure and flow data. Fourier transforms were then performed on this data to calculate $Z_0$ and $Z_1$. Fiber bundle resistance was calculated by averaging bundle pressures:

$$R = \frac{\left(\frac{P_3 + P_5}{2}\right) - \left(\frac{P_4 + P_6}{2}\right)}{Q} \quad (15)$$

Viscous resistance was calculated at $Q=4$ L/min and $HR=100$ beats/min for the inlet, bundle and outlet section of each model in a similar fashion as the CFD simulation. Stroke volume, $SV$, was calculated as $Q/HR$. Correlations between $Z_0$ and $SV$ and between $Z_1$ and $Q$ were created.

Comparisons between \textit{in vitro} $Z_0$ and $Z_1$ were performed with SPSS (SPSS, Chicago, IL) using a mixed model with housing prototype number as the subject variable, stroke volume as a fixed, repeated-measure variable, and $\theta$ as a fixed variable.
Results

CFD: Rigid Housing TAL

Figure 2.5 displays TAL $Z_0$ CFD results for varying stroke volumes at a pulsatility of 3.75.

![Graph showing TAL Z0 for varying stroke volumes](image)

Figure 2.5: Effect of $\theta$ on TAL $Z_0$ for varying stroke volumes: comparison between in vitro and CFD results

Device $Z_0$ increases linearly as $SV$ increases, as well as increasing with larger $\theta$. At $Q=4$ L/min and $HR=100$ beats/min ($SV=0.04$ L), $Z_0$ was 0.47, 0.61, and 0.79 mmHg/(L/min) for the $15^\circ$, $45^\circ$, and $90^\circ$ devices, respectively. The correlations relating $Z_0$ to $SV$ were:

$$Z_0 = 1.82SV + 0.41, \ R^2=97.7\%, \ for \ \theta=15^\circ$$  \hspace{1cm}(16)

$$Z_0 = 6.17SV + 0.39, \ R^2=97.4\%, \ for \ \theta=45^\circ$$  \hspace{1cm}(17)

$$Z_0 = 10.7SV + 0.36, \ R^2=98.7\%, \ for \ \theta=90^\circ$$  \hspace{1cm}(18)
Figure 2.6 displays $Z_I$ CFD results at varying heart rates and flow rates at a pulsatility of 3.75.

The values of $Z_I$ increase linearly with flow rate for the 15° model; however, they remain more uniform at each flow rate for the 45° and 90° models. $Z_I$ is larger at $HR=100$ beats/min for all three models. Lastly, $Z_I$ was similar between the 45° and 90° models but noticeably greater for the 15° model. At $Q=4$ L/min and $HR=100$ beats/min, $Z_I$ was 0.67, 0.40, and 0.35 mmHg/(L/min) for the 15°, 45°, and 90° models, respectively.

Device $Z_0$ was reduced slightly under more typical in vivo conditions with a lower inlet flow pulsatility; however, $Z_I$ was not affected. At $Q=4$ L/min and $HR=100$ beats/min, the 45° model $Z_0$ was 0.53 mmHg/(L/min) at a pulsatility of 2 vs. 0.61 mmHg/(L/min) at a pulsatility of 3.75. The 45° model $Z_I$ was 0.41 mmHg/(L/min) at a pulsatility of 2 vs. 0.40 mmHg/(L/min) at a pulsatility of 3.75. Lastly, fiber bundle
resistance for all models was 0.31 mmHg/(L/min). The majority of device resistance came from the fiber bundle and inlet housing sections in each design. Table 2.1 lists the viscous resistances in the inlet, bundle, and outlet sections for the 45° and 90° models at an example condition of $Q=4$ L/min and $HR=100$ beats/min.

Table 2.1: Viscous Resistance in the inlet, bundle and outlet sections for the 45° and 90° in vitro and CFD models at $Q=4$ L/min and $HR=100$ beats/min.

<table>
<thead>
<tr>
<th>Resistance</th>
<th>In Vitro, 45°</th>
<th>CFD, 45°</th>
<th>In Vitro 90°</th>
<th>CFD 90°</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inlet, mmHg/(L/min)</td>
<td>0.25 ± 0.06</td>
<td>0.21</td>
<td>0.26 ± 0.09</td>
<td>0.31</td>
</tr>
<tr>
<td>Inlet, % of Total</td>
<td>36.2 ± 10.5%</td>
<td>34.2%</td>
<td>32.2 ± 10.9%</td>
<td>39.4%</td>
</tr>
<tr>
<td>Bundle, mmHg/(L/min)</td>
<td>0.31 ± 0.11</td>
<td>0.31</td>
<td>0.28 ± 0.09</td>
<td>0.31</td>
</tr>
<tr>
<td>Bundle, % of Total</td>
<td>43.5 ± 14.1%</td>
<td>49.4%</td>
<td>35.1 ± 12.9%</td>
<td>38.7%</td>
</tr>
<tr>
<td>Outlet, mmHg/(L/min)</td>
<td>0.14 ± 0.16</td>
<td>0.10</td>
<td>0.26 ± 0.12</td>
<td>0.17</td>
</tr>
<tr>
<td>Outlet, % of Total</td>
<td>20.3 ± 19.2%</td>
<td>16.4%</td>
<td>32.7 ± 12.4%</td>
<td>22.0%</td>
</tr>
<tr>
<td>Total, mmHg/(L/min)</td>
<td>0.70 ± 0.14</td>
<td>0.62</td>
<td>0.81 ± 0.10</td>
<td>0.79</td>
</tr>
</tbody>
</table>

Band plots taken at peak systole for $Q = 4$ L/min, $HR = 100$ beats/min, and pulsatility = 3.75 and 2 are shown in Figure 2.7(a) along a section in the device inlet region for two different plot ranges. These plots indicate areas of little to no flow at the distal end of the 15° and 45° device, as indicated by the blue color. Also, the wider range (0-0.014 m/s) plots indicate that flow is more even in the inlet housing of the 15° model. A smaller pulsatility of 2 seems to result in slightly more even flow, but with large low flow regions. Band plots taken during systole across the bundle indicate uniform flow in
the 15° and 45° model with slightly lower flow near the front of the device in the 90° model (Figure 2.7(b)).

During diastole (Figure 2.8), no more flow is entering the device and velocities are lower overall; however, velocities are higher in the 45° and 90° models due to recirculation. The extent of this recirculation is more easily visualized with velocity vector plots.

Figure 2.7: Velocity band plots for the 15° and 90° CFD TAL models at a pulsatility of 3.75 and the 45° model at a pulsatility of 3.75 and 2. Shown at peak systole in the (a) inlet housing region and (b) fiber bundle.
Velocity vector plots of the inlet housing region, taken at the beginning of diastole (t=1.58s) for $Q = 4$ L/min, $HR = 100$ beats/min, and pulsatility = 3.75, are shown in Figure 2.9.

Figure 2.8: Velocity band plots for the 15° and 90° CFD TAL models at a pulsatility of 3.75 and the 45° model at a pulsatility of 3.75 and 2. Shown at end-diastole for two different velocity plot ranges.
A magnified view of the major recirculation regions (indicated in Figure 2.9(a)) are shown in Figure 2.9(b). Large vortices occur near the distal end of the 90° device while smaller recirculation regions occur in the proximal end of the 15° and 45° devices. In the 90° model at peak systole (Figure 2.10), large vortices occur in the proximal end, as well as smaller one at the distal end of the device.
The large vortices in this device travel from the proximal end to the distal end of the model (with flow) as time in the cardiac cycle progresses (Figure 2.11). The 15° and 45° models also have vortices occurring throughout the cardiac cycle; however, these are smaller and mainly occur in the proximal end of the device. Overall, a lower pulsatility of 2 results in less recirculation in the housing; however, like the 15° model, there is a large low flow region at the distal end of the device (Figure 2.11).
Velocity vector plots taken in the housing outlet (Figure 2.12) showed small regions of recirculation in the outlet tube (Figure 2.12b) and downstream end of the housing (Figure 2.12c) during diastole only. However, the velocities in these vortices were very small compared to those in the housing inlet.

Figure 2.11: Velocity vector plots for the 15° and 90° CFD TAL models at a pulsatility of 3.75 and the 45° model at a pulsatility of 3.75 and 2. Shown at end-diastole in the inlet housing region.
In Vitro Testing

The effect of $\theta$ on TAL in vitro $Z_0$ and $Z_1$ is shown in Figures 2.5 and 2.13, respectively, for an average pulsatility of 3.3. $Z_0$ varied significantly with $SV$ ($p<10^{-19}$) and $\theta$ ($p<0.01$). The 45° model had a significantly lower $Z_0$ than the 90° model overall. At $Q=4$ L/min and $HR=100$ beats/min ($SV=0.04$ L), $Z_0 = 0.69\pm0.13$ and $0.79\pm0.10$ mmHg/(L/min) for the 45° and 90° models, respectively. The correlations relating in vitro $Z_0$ to $SV$ (in L) were:

$$Z_0 = 8.889SV + 0.297, \ R^2=91.1\%, \text{ for } \theta=45^\circ \ (19)$$

$$Z_0 = 9.084SV + 0.396, \ R^2=93.0\%, \text{ for } \theta=90^\circ \ (20)$$

Figure 2.12: (a) Velocity vector plot for the 45° TAL model at the beginning of diastole ($t=1.58s$) in the outlet housing region. Magnified view of the outlet tube (b) and the proximal end of the housing (c) showing small recirculation regions.
Values for $Z_1$ varied significantly with $HR$ ($p<0.01$), $Q$ ($p<10^{-30}$), and $\theta$ ($p<10^{-6}$). The effect of $HR$ was very small compared to $Q$ and $\theta$, but indicated that $Z_1$ increased with $HR$. The 90° model had significantly lower $Z_1$ than the 45° model. At $Q=4\text{ L/min}$ and $HR=100\text{ beats/min}$, $Z_1 = 1.23\pm0.27$ and $0.91\pm0.0.13\text{ mmHg/(L/min)}$ for the 45° and 90° models, respectively. The correlations relating in vitro $Z_1$ for each $\theta$ and $HR$ are:

$$Z_1 = 0.096Q + 0.737, \ R^2=99.4\%, \text{ for } \theta=45^\circ, \ HR=80\text{ beats/min} \quad (21)$$

$$Z_1 = 0.118Q + 0.789, \ R^2=98.6\%, \text{ for } \theta=45^\circ, \ HR=100\text{ beats/min} \quad (22)$$

$$Z_1 = 0.165Q + 0.222, \ R^2=99.99\%, \text{ for } \theta=90^\circ, \ HR=80\text{ beats/min} \quad (23)$$

$$Z_1 = 0.189Q + 0.192, \ R^2=99.4\%, \text{ for } \theta=90^\circ, \ HR=100\text{ beats/min} \quad (24)$$

Figure 2.13: Effect of $\theta$ on TAL $Z_1$: comparison between in vitro and CFD results for varying heart and flow rates
Overall, *in vitro* $Z_1$ were larger and showed a stronger dependence on $\theta$ and $Q$ than CFD results. These larger $Z_1$ were due to larger peak pressures *in vitro*, even though average pressure drops were similar between the CFD and *in vitro* studies.

Average fiber bundle resistance for all models was 0.29 mmHg/(L/min), nearly identical to CFD results. In the 45° model, the majority of device resistance came from the fiber bundle and inlet housing sections. However, in the 90° model, the relative contribution to total device resistance was almost identical for each section (Table 2.1).

Lastly, at a lower pulsatility of 1.7 with $Q=4$ L/min and $HR=100$ beats/min ($SV=0.04$ L), the 45° model had $Z_0=0.65$ mmHg/(L/min) and $Z_1=1.29$ mmHg/(L/min). The $Z_0$ and $Z_1$ are 0.69 and 1.23 mmHg/(L/min) at the same conditions with pulsatility equal to 3.2. Therefore, as predicted in the CFD study, pulsatility does not greatly influence impedance *in vitro*.

**Discussion**

The impedances of all the TAL designs tested in this study are significantly lower than previous designs. Experiments with the MC³ Biolung (MC³, Ann Arbor, MI) [9] and a cTAL [3] report $Z_0=1.8$ and 1.9 mmHg/(L/min), respectively at 3.0 cP glycerol flows of $Q=4$ L/min and $HR=100$ beats/min. At the same flow and heart rate, the 45° and 90° model in the current study had $Z_0=0.69\pm0.13$ and $0.79\pm0.10$ mmHg/(L/min). These values are also significantly lower than the impedance of the natural lung: $Z_0=1.25$ mmHg/(L/min). $Z_1$ in the healthy, natural lung is 0.49 mmHg/(L/min) [10] compared to 1.23 mmHg/(L/min) in the 45° model in this study.
The effect of $SV$ and $\theta$ on $Z_0$ was very similar in the CFD and in vitro results. However, CFD modeling predicted lower $Z_1$ values overall with a minimal effect of $\theta$ and $Q$. The $Z_1$ is more heavily influenced by differences in flow rate patterns and thus has a higher variability than $Z_0$ during in vitro testing. Overall, the $Z_1$ values measured for both models are small and unlikely to influence device function in vivo [11].

The 45° and 90° devices tested in this study had a 62 and 56% lower in vitro $Z_0$ than previous designs. This decrease should improve RV function during attachment in vivo. The ultimate effect of lower impedance on cardiac function depends on attachment mode. The most likely clinical application for these devices is attachment in parallel with the natural lung in a PA to left atrium (LA) configuration. If attached in parallel with the natural lung, lower TAL impedance may allow patients to more easily exercise during device attachment. Akay et al attached the MC3 Biolung in parallel with the pulmonary circulation in sheep with chronic pulmonary hypertension and varied the percentage of CO through the device under simulated rest, ambulatory, and exercise conditions [1]. At ambulatory and resting conditions, CO was maintained as the percentage of CO to the device increased. At exercise conditions, CO decreased with increasing flow to the TAL up to $23 \pm 5\%$ at 90% flow diverted through the TAL. TAL impedance increases and the natural lung impedance decreases with increasing blood flow. Thus, during exercise with elevated CO, a higher percentage of CO may travel through the native lungs reducing respiratory support when attached in parallel. Further decreases in device impedance may allow for complete exercise tolerance and optimal device performance.

If attached in series with the natural lung, from the proximal to distal PA, lower impedance should reduce the drop in CO that has been observed experimentally.
Previous studies by Kim et al, Kuo et al, and Akay et al developed relationships between acute increases in $Z_0$ and $Z_1$ and the resultant percent decrease in CO [11-13]. These studies can be used to predict the improvement in CO using these devices vs. previous designs. In the devices tested here, $Z_1$ values are small in all devices, regardless of design, and should not have a measureable effect on CO [11]. Values of $Z_0$ should, on the other hand, contribute to lowering CO. The increase in pulmonary system impedance ($\Delta Z$) due to the TAL impedance ($Z_{TAL}$) can be calculated with the equation:

$$\Delta Z = Z_{TAL} f$$  \hspace{1cm} (25)

where $f$ is the fraction of flow to the TAL. If we assume CO to be 6 L/min and allow 4 L/min to travel through the TAL then $f=2/3$. Using the 45° model in vitro data at $Q=4$ L/min, $HR=100$ beats/min, pulsatility=1.7, and adding 0.87 mmHg/(min) for the anastomoses resistance [13], $\Delta Z_0$ and $\Delta Z_1$ are 1.01 and 0.86 mmHg/(L/min), respectively. The Kim, Kuo, and Akay relationships predict a 4.1, 13, and 7.6% decrease in CO, respectively. For the previous TAL designs, we would predict a decrease of 15, 19, and 13%. Based on these predictions, the 45° model should cause a small decrease in CO. If we average the Kim, Kuo, and Akay data, $f=0.75$ will give a 10% reduction in CO. This decrease may be tolerable in patients with normal pulmonary vascular resistance or mild pulmonary hypertension and normal right ventricular function. It would not in patients with marked pulmonary hypertension and right ventricular dysfunction. If necessary, $f$ could be reduced to further decrease the drop in CO.

The CFD model also provided important information on how device design affects flow patterns in both the housing and fiber bundle. Changes in $Z_0$ can be explained in large part by examining blood flow recirculations in the TAL. Differences in flow
patterns were more pronounced in the housing inlet. The fiber bundle acted to evenly distribute blood flow, resulting in more even blood flow in the housing outlet and only small, low-velocity, recirculations. Recirculation occurred in the inlet of all three models; however, recirculations decreased as $\theta$ decreased. The 90° housing inlet had several regions of large vortices throughout the cardiac cycle, while the 15° and 45° models had smaller areas of recirculation.

Device design also affected flow uniformity in the housing inlet and fiber bundle. Flow uniformity is essential to the short and long-term function of the device. Regions of little or no flow offer little gas exchange when they occur in the fiber bundle and, worse yet, can lead to a buildup of activated clotting factors and clot formation when they occur at any of the device artificial surfaces. The 15° and 45° models had areas of little to no flow near the distal end of the inlet. These areas were larger in the 15° model where flow was barely reaching past the middle of the housing body. The stasis regions in the back of the device can never completely be eliminated for the angled housings, but further geometry changes may help, such as a shorter distance between the inlet and the back and tapered corners in the back. Tapered corners are easily incorporated during the production process when the fiber bundle is “potted” in urethane to separate the blood and gas phases of the device.

Ultimately, the optimal device must have a combination of low impedance, even flow, and reasonable size for use by an ambulatory patient. While the 15° model had smaller recirculation regions and low impedance, it had a large stasis region near the distal end of the device. Also, the long inlet and outlet length of the 15° model may make it unwieldy to use with patients. Therefore, the 45° model has the optimal combination
of low impedance, better flow patterns, and relatively compact size. The low impedance of the 45° model should allow for a minimal, acute decrease in CO at rest if implanted in series with the natural lung and increased exercise tolerance when compared to previous TAL designs if attached in parallel with high flows to the TAL. These results will be confirmed by examining compliant housing models, discussed in the following chapters.
References


Introduction

CFD and in vitro studies with a rigid-housing TAL showed that device impedance was reduced as the inlet and outlet expansion angle, \( \theta \), decreased. The optimum combination of low impedance and good flow patterns was achieved with the 45° housing model. Now we will confirm these results and investigate whether additional reductions in TAL impedance may be achieved by using a compliant housing material. Unlike a hard-shell TAL, the compliant housing can expand to accommodate the large changes in blood volume seen during a heartbeat. This dampens the flow pulse, creating a more steady flow through the cardiac cycle.

This study aimed to reduce TAL impedance using a compliant housing material in conjunction with more gradual inlet and outlet geometries, as explored in Chapter 2. In this compliant thoracic artificial lung (cTAL), blood flows through the inlet and into the compliant inlet manifold (Figure 2.1). The housing expands, accommodating the stroke volume without a large pressure increase. Blood then flows through the fiber bundle where gas exchange occurs, through the bottom housing manifold and finally exits through the outlet. The effect of inlet and outlet expansion angle, \( \theta \), on cTAL impedance and flow patterns was explored with fluid-structure interaction (FSI) analysis. The FSI
analysis was necessary to examine the interaction between fluid forces and housing deformation. The optimal cTAL design was determined from FSI results, prototyped, and tested in vitro to assess impedance and gas transfer performance.

Methods

Fluid Structure Interaction Analysis

SolidWorks was used to create cTAL models with varying housing geometries using $\theta = 15^\circ$, $45^\circ$, and $90^\circ$ (Figures 2.1 and 2.2). The inlet/outlet diameter, height, and width of each cTAL model are 0.016 m, 0.103 m, and 0.102 m, respectively. The fiber bundle length, path length, and frontal area are 0.127 m, 0.038 m, and 0.013 m$^2$, respectively, for each model. The $15^\circ$, $45^\circ$, and $90^\circ$ models have a total length of 0.338, 0.221 m, and 0.178 m. These models were then imported into ADINA. For FSI analysis, separate structure and fluid models were created, coupled and then run together. The fluid model consisted of all regions that fluid passed through while the structure model consisted of the Biospan housing.

FSI: Problem Formulation

Program assumptions included a transient solution with incompressible flow. At a flow of 3 L/min, the Reynolds number during peak flow was 4150 in the inlet section, 170 coming out of the bundle into the outlet section, and 3300 exiting the device. Due to limitations in the CFD software, we could not apply turbulent flow to the fiber bundle (porous media model). This required us to split up sections and run their simulations
independently, while carrying boundary conditions from one section to the next. Two such solutions were attempted: i) running each section independently, with turbulence in the inlet and outlet sections and laminar flow in the bundle (as done for TAL models in Chapter 2), and ii) running the inlet section as turbulent and remainder of the model as laminar. The former gave unrealistic flow patterns and dampening (detailed further in discussion). The latter gave us the most quantitatively accurate and physically realistic results possible and is thus presented here.

In the fluid model, the properties of the fiber bundle and the fluid characteristics of blood were defined as described in the TAL CFD study (Chapter 2). A slip wall boundary condition was applied to the bundle walls while a FSI condition was used in the inlet and outlet regions. A pressure of 770 mmHg (absolute pressure) was applied at the outlet and also used for the initial pressure condition. A time dependent velocity was applied to the inlet face using the same pulsatile flow waveform (as described in Chapter 2) with average flow rates, $Q$, of 3 and 5 L/min and a heart rate, $HR$, of 100 beats/min.

The structure model consisted of the compliant, Biospan (DSM PTG, Berkeley, CA) housing, with a thickness of 0.51 mm, forming the inlet and outlet sections of the model. Biospan was defined as an elastic isotropic material with a Young’s modulus of 6.21 MPa, Poisson’s ratio of 0.40, and density of 1030 kg/m$^3$. FSI boundary conditions were applied to each face and shell elements were used with a face thickness of 0.51 mm. The edges bordering the fiber bundle were fixed in all directions, along with the inlet and outlet edge to prevent translation of the model in space. Symmetry plane edges were fixed in the direction normal to the symmetry plane. Atmospheric pressure was applied to
the model by applying a pressure load of 760 mmHg to the exterior faces of the structure model.

The governing equations used for the problem formulation in the fluid domain are the continuity and Navier-Stokes equations:

\[ \nabla \cdot \mathbf{u} = 0 \]
\[ \rho_f \frac{\partial \mathbf{u}}{\partial t} + \rho_f (\mathbf{u} - \mathbf{u}_g) \cdot \nabla \mathbf{u} = \nabla \cdot \mathbf{\sigma}_f + \rho_f \mathbf{f}_f^B \]  \hspace{1cm} (1)

where \( \mathbf{u} \) is the fluid velocity vector, \( \rho_f \) is the fluid density, \( \mathbf{u}_g \) is the moving coordinate velocity, \( \mathbf{f}_f^B \) is the body force per unit volume, \( (\mathbf{u} - \mathbf{u}_g) \) is the relative velocity of the fluid with respect to the moving coordinate velocity, and \( \mathbf{\sigma}_f \) is the fluid stress tensor.

The governing equation for the solid domain of the FSI model is the elastodynamics equation:

\[ \rho_s \ddot{\mathbf{d}}_s = \nabla \cdot \mathbf{\sigma}_s + \mathbf{f}_s^B \]  \hspace{1cm} (2)

where \( \rho_s \) is the wall density, \( \mathbf{\sigma}_s \) is the solid Cauchy stress tensor, \( \mathbf{f}_s^B \) is the externally applied body force vector at time \( t \) and \( \ddot{\mathbf{d}}_s \) represents the local acceleration of the solid region (\( \ddot{\mathbf{d}}_s = \ddot{\mathbf{u}}_g \)).

The boundary conditions at the interface between the fluid domain and the solid domain are:

Displacement : \( \mathbf{d}_f = \mathbf{d}_s \)

Traction : \( \mathbf{\sigma}_f = \mathbf{\sigma}_s \)

No-slip : \( \mathbf{u} = \mathbf{u}_g \) \hspace{1cm} (3)

where \( \mathbf{d}, \mathbf{\sigma}, \mathbf{n} \) are the displacement vector, stress tensor, and normal vector, respectively.

The subscripts \( s \) and \( f \) indicate fluid and solid properties, respectively.
FSI: Numerical Scheme

A finite element formulation based on the Galerkin method (described in Chapter 2) was utilized to solve the governing equations. Direct computing of two-way fluid-structure coupling was used. Convergence criteria were set at $\leq 1.5\%$ difference in impedance between models of different time step or mesh size. The final mesh size was 3 mm for the bodies, 0.25 mm for the inlet and outlet sections, and 1.0 mm for the inlet and outlet faces. A time step of 0.001 s was used.

FSI: Data Analysis

The fluid and structure models were run simultaneously with FSI analysis for four consecutive heart beats to achieve a converged, periodic solution. Inlet and outlet pressure data, along with flow, from the final beat were obtained from the program. $Z_0$ and $Z_I$ were calculated and used to assess cTAL impedance. Pressure and flow data was also obtained from the top and bottom fiber bundle faces to determine bundle resistance. Average maximum housing displacement, $D_{\text{avg}}$, was also calculated, from absolute magnitude values, over the last heartbeat for the inlet and outlet manifold of each model at each flow condition.

ADINA was also used to create velocity vector plots, at the beginning of diastole in the cardiac cycle, to investigate areas of recirculation in each housing model. In these plots, color and the size of the vectors are used to indicate velocity magnitude where red represents the maximum velocity in the plot range and blue represents a velocity of 0 m/s. Velocity band plots were also generated to assess flow uniformity in the housing and
bundle of each model. Plots were created on a cross section along the y-axis (top down view) for cross sections at the entrance to the fiber bundle from the inlet section and in the middle of the fiber bundle. In these plots, color denotes velocity magnitude with red representing the highest velocity in the plot range and blue representing a velocity of 0 m/s. A plot range of 0-0.0081 m/s was created at peak systole in the cardiac cycle.

**Compliant Thoracic Artificial Lung**

The cTAL model with \( \theta = 45^\circ \) was prototyped for *in vitro* testing (Figure 3.1).

![Figure 3.1: cTAL prototype showing blood flow path](image)

The compliant housing was created by dipping a mold into Biospan (DSM PTG, Berkeley, CA), a polyurethane polymer, until a housing thickness of 0.51 mm was achieved. The device also consisted of a bundle of polypropylene hollow fibers which had a fiber diameter, porosity, path length, frontal area, and surface area of 210 \( \mu \)m, 0.75, 0.038 m, 0.013 m\(^2\), and 2.4 m\(^2\), respectively. Device compliance was determined using previously reported methods [1]. Fluid volume was slowly added to the device, while
pressure was recorded and compliance was calculated as $\Delta V/\Delta P$, until an added fluid volume of 560 mL was reached.

*In Vitro Testing: Hemodynamics*

*In vitro* testing was conducted in a circuit consisting of a pulsatile pump (Harvard Apparatus, Holliston, MA), cTAL ($n = 4$), and reservoir filled with 3.0 cP glycerol solution (Figure 3.2a). The reservoir was raised 12.6 cm above the cTAL outlet, resulting in a constant device outlet pressure of 10 mmHg. Fluid-coupled pressure transducers (Abbott Critical Care Systems, Chicago, IL) measured pressure at the cTAL inlet ($P_i$) and outlet ($P_o$). A flow probe (14PXL, Transonic Systems Inc, Ithaca, NY) measured the flow rate into the cTAL ($Q_{cTAL}$).
The pulsatile pump created six flow conditions: heart rates (HR) of 80 and 100 beats/min and flow rates (Q) of 2, 4, and 6 L/min. At each condition, a data set of $Q_{CTAL}$, $P_i$, and $P_o$ was digitally acquired for 15 seconds at a sampling frequency of 250 Hz through a BIOPAC data acquisition system (BIOPAC, Goleta, CA). Fourier transforms were then performed on this data to calculate $Z_0$ and $Z_1$. Stroke volume, $SV$, was calculated as $Q/HR$, and linear regressions were performed between $Z_0$ and $Z_1$ and $SV$.

Figure 3.2: (a) In vitro testing circuit for resistance study consisting of pulsatile pump, cTAL, and reservoir filled with 3.0 cP glycerol solution. Two pressure lines were connected to the model at the inlet (P1) and outlet (P2). (b) Gas transfer testing circuit consisting of a reservoir filled with bovine blood, roller pump, deoxygenator (conditioning loop) and a cTAL (testing loop).
In Vitro Testing: Gas Transfer

The gas transfer performance of four cTALs was tested in a recirculation circuit consisting of bovine blood (Figure 3.2b). Blood was pumped by a roller pump (Baxter Century, Baxter, Deerfield, IL) from the recirculation reservoir, through a deoxygenator (Capiox Rx 25, Terumo Cardiovascular Systems, Ann Arbor, MI) and returned to the reservoir in order to bring the blood to venous conditions according to AAMI protocols [2]. The sweep gas used to condition the blood consisted of varying levels of O2, N2, and CO2. Once the blood was conditioned, blood flow was routed from the pump to the cTAL. Blood was pumped through the cTAL at flow rates of 3, 5, and 7 L/min, and oxygen flow was set at a gas to blood flow ratio of two.

Blood samples were taken from the cTAL inlet and outlet and analyzed to determine PO2, PCO2, pH, SO2, and hemoglobin concentration. The O2 and CO2 transfer rates were calculated with the following standard formulas:

\[
VO_2 = Q_b (k(p_o-p_i)+1.34c_{hb}(s_o-s_i)) \quad (4)
\]

\[
VCO_2 = Q_g c_{CO2} \quad (5)
\]

where \(Q_b\) is the blood flow rate, \(k\) is the solubility of oxygen in blood, \(p_o\) is outlet PO2, \(p_i\) is inlet PO2, \(c_{hb}\) is the hemoglobin concentration, \(s_o\) is the outlet fractional oxyhemoglobin saturation, \(s_i\) is the inlet fractional oxyhemoglobin saturation, \(Q_g\) is the sweep gas flow rate, and \(c_{CO2}\) is the exit sweep gas carbon dioxide fraction.
Results

Fluid Structure Interaction Analysis

The compliant housing of the cTAL caused dampening of device outlet flow. Inlet and outlet flow waveforms for $Q=3$ L/min are shown in Figure 3.3.

Figure 3.3: cTAL inlet and outlet flow waveforms for the FSI models at 3 L/min and 100 beats/min, showing outlet flow dampening. Representative cTAL inlet flow waveform from the in vitro study at 4 L/min and 100 beats/min.

Figure 3.4 displays FSI results with cTAL $Z_0$ and $Z_1$ on the y axis and $SV$ on the x axis.
For the 15° and 45° models, $Z_0$ increases minimally as $SV$ increases. In the 90° model, however, $Z_0$ increased 13% from $SV=0.03$ to 0.05 L. At $SV=0.03$ L, $Z_0$ was 0.50, 0.69, and 0.92 mmHg/(L/min) for the 15°, 45°, and 90° devices, respectively. $Z_1$ increases with increasing $\theta$ and at $SV=0.03$ L, $Z_1$ is 0.16, 0.50, and 0.93 mmHg/(L/min) for the 15°, 45°, and 90° models, respectively. Velocity vector plots in the inlet region (Figure 3.5) depict recirculation regions in the inlet that increase in size and velocity with $\theta$, explaining the effect of $\theta$ on $Z_0$.

Figure 3.4: FSI results showing the effect of $\theta$ on cTAL $Z_0$ and $Z_1$ for varying stroke volumes
Velocity band plots taken at cross sections in the inlet region at the entrance to the fiber bundle (Figure 3.6a) and the middle of the fiber bundle (Figure 3.6b) are shown in Figure 3.6.
Figure 3.6a shows areas of low flow in the distal end of the housing in the 15° and 45°
models. At the entrance to the fiber bundle, flow is mostly uniform at the proximal end of
all models, except for a small spot in the 45° and 90° models. Figure 3.6b shows uniform
flow in the fiber bundle, with very small velocity differences between the proximal and
distal ends. Velocity in the bundle at peak systole in the 90° model is slightly higher than
the other models. This is due to less dampening of the flow pulse in the inlet region of the
90° model, causing more blood to enter the fiber bundle during systole (see below).

Figure 3.7 shows the flow waveforms entering the fiber bundle for each model. Peak flow
is highest for the 90° model, while flow grows more even across the cardiac cycle with
decreasing θ.
Table 3.1: Maximum housing displacement in one heart beat, $D_{\text{max}}$, and average maximum housing displacement over one heart beat, $D_{\text{avg}}$, for the inlet and outlet housing sections for each model and flow rate.

<table>
<thead>
<tr>
<th>Model, $\theta$</th>
<th>Flow Rate, Q, L/min</th>
<th>Inlet $D_{\text{avg}}$ (mm)</th>
<th>Outlet $D_{\text{avg}}$ (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15°</td>
<td>3</td>
<td>1.97</td>
<td>1.82</td>
</tr>
<tr>
<td>15°</td>
<td>5</td>
<td>2.29</td>
<td>2.04</td>
</tr>
<tr>
<td>45°</td>
<td>3</td>
<td>1.8</td>
<td>1.66</td>
</tr>
<tr>
<td>45°</td>
<td>5</td>
<td>2.08</td>
<td>1.86</td>
</tr>
<tr>
<td>90°</td>
<td>3</td>
<td>1.64</td>
<td>1.51</td>
</tr>
<tr>
<td>90°</td>
<td>5</td>
<td>1.95</td>
<td>1.73</td>
</tr>
</tbody>
</table>

Figure 3.7: Flow waveforms entering the fiber bundle for the 15°, 45°, and 90° cTAL models
Flow dampening occurs due to the housing expanding and contracting with the flow pulse. The average maximum housing displacement over one heart beat, $D_{avg}$, for the inlet and outlet housing sections is given in Table 3.1 for each model and flow rate. With increasing $Q$, $D_{avg}$ increases for both housing sections. Housing displacement is larger in the inlet section than in the outlet section. Displacement occurs more gradually over the flow pulse for smaller $\theta$, causing $D_{avg}$ to be largest in the 15° model.

In Vitro Testing: Hemodynamics

The cTAL had two different compliance values, which were dependent on pressure. At low pressures, the device was still filling with fluid and the housing hadn’t begun stretching, resulting in a large compliance. At higher pressures, the housing of the cTAL stretched to accommodate the increase in fluid, and the compliance was smaller. Figure 3.8 shows the compliance curve for a typical device. Average cTAL compliance was 93.9 ± 8.75 mL/mmHg for internal pressure < 5 mmHg and 5.21 ± 0.57 mL/mmHg for internal pressure > 5 mmHg.
In vitro $Z_0$ and $Z_1$ data for varying $SV$ are displayed in Figure 3.9.

Figure 3.8: *In vitro* compliance curve for a typical device. The slope of the volume vs. pressure linear curves at high pressure and low pressure is equal to the compliance in each corresponding pressure zone.
As SV increases, $Z_0$ decreases slightly from $0.53 \pm 0.08 \text{ mmHg/(L/min)}$ at $SV=0.02 \text{ L}$ to $0.41 \pm 0.14 \text{ mmHg/(L/min)}$ at $SV=0.075 \text{ L}$. The correlation relating $Z_0$ to $SV$ was: $Z_0 = -2.691SV + 0.615$, $R^2=65\%$. The value of $Z_1$ increases with $SV$, with a correlation of $Z_1 = 9.472SV + 0.485$, $R^2=75\%$. In vitro, $Z_1$ values are higher than the FSI models predicted, while experimental $Z_0$ values are slightly lower.

In Vitro Testing: Gas Transfer

Gas transfer performance is shown in Table 3.2.
Table 3.2: cTAL gas transfer performance: outlet PO2, outlet SO2, VO2 and VCO2 for varying blood flow rates.

<table>
<thead>
<tr>
<th>Blood Flow, Q, L/min</th>
<th>Outlet PO2 (mmHg)</th>
<th>Outlet SO2 (%)</th>
<th>VO2 (mL/min)</th>
<th>VCO2 (mL/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>436.0 ± 15.9</td>
<td>99.9 ± 0.02</td>
<td>196.6 ± 13.4</td>
<td>330.0 ± 73.2</td>
</tr>
<tr>
<td>5</td>
<td>360.9 ± 54.4</td>
<td>99.9 ± 0.04</td>
<td>307.6 ± 18.0</td>
<td>417.5 ± 91.8</td>
</tr>
<tr>
<td>7</td>
<td>242.8 ± 48.9</td>
<td>99.7 ± 0.14</td>
<td>395.4 ± 42.9</td>
<td>500.5 ± 35.0</td>
</tr>
</tbody>
</table>

The outlet PO2 decreases with blood flow rate but remains above 240 mmHg at all conditions. Similarly, SO2 remains above 99.5%. Lastly, VO2 and VCO2 increase linearly with increasing blood flow rate.

**Discussion**

FSI impedance and flow patterns were used to choose the optimal model for prototyping. As $\theta$ decreased, impedance decreased. At $Q$= 3 L/min and $HR$= 100 beats/min, $Z_0$ values for the 15°, 45°, and 90° models were 0.51, 0.69, and 0.92 mmHg/(L/min), respectively. $Z_1$ values were low, under 1 mmHg/(L/min), for all three models. For the range of $Z_0$ values found here, an increase in $Z_1$ of 12-24 mmHg/(L/min) is necessary in order to decrease CO by 1% [3]. Thus, the $Z_1$ differences found in this study are inconsequential, would have no significant effect on cardiac function, and thus have no bearing on device selection.

Based on these results alone, $\theta$ = 15° would appear to be the ideal model. However, in the 15° model, blood flow enters with such a limited velocity that it does not
reach the back end of the manifold, creating a large low flow region at the distal end of the device. This low flow region could lead to reduced gas transfer performance and clot formation over time. The 45° device had a smaller, low flow region at the distal end. The 90° model possesses a higher velocity inlet jet that reaches the back end of the device, leading to largely uniform flow. Flow through the bundle was mostly uniform in all models, with slightly higher velocities at the proximal end. While flow was more spatially uniform with higher $\theta$, flow was also less temporally uniform due to lower dampening of the flow pulse. Low flow during diastole could also lead to stasis and clot formation.

The 15° model has low impedance and more even flow patterns throughout the cardiac cycle but poor flow at the distal end of the device. Moreover, the 15° model is excessively long, giving it a large priming volume and making it impractical for paracorporeal use. The 90° model has good flow uniformity at the distal end of the device but a period of low flow during systole and higher impedance. Thus, the 45° model was chosen for prototyping and \textit{in vitro} testing due to its combination of low impedance, good flow patterns, and ease of manufacture.

\textit{In vitro} testing demonstrated similar, yet lower values for $Z_0$ than FSI modeling. The FSI model was unable to completely mirror the experimental conditions. Due to limitations in the CFD software, we were required to split up the model and run the inlet section as turbulent and remainder of the model as laminar. In the TAL CFD models (Chapter 2) we were able to run both the inlet and outlet sections as turbulent, giving results that matched \textit{in vitro} impedance values much more closely. However, when attempting this using FSI analysis, unrealistic flow profiles were seen in the outlet
housing section. This gave more quantitatively accurate results for $Z_0$, but unrealistic flow patterns. Our final FSI flow assumptions gave us relatively good accuracy for both; however, being able to apply turbulent flow uniformly throughout the model may have resulted in a more accurate CFD model, resulting in lower $Z_0$ values. Regardless of the quantitative results, qualitative results were similar and the 45° model was chosen as the optimal design.

Like the rigid-housing results, the cTAL impedance found with this design is much lower than all previous TAL designs and, for that matter, than a healthy native lung itself. Previous experiments examining the impedance of the MC³ Biolung (MC³, Ann Arbor, MI) [4] and an early cTAL [1] reported $Z_0=1.8$ and 1.9 mmHg/(L/min), respectively at $Q=4$ L/min and $HR=100$ beats/min. Studies with the latter device suggested the majority of device resistance was from the inlet and outlet regions. The low resistance in the cTAL in the current study is a function of three things: a gradual low resistance inlet, a low resistance fiber bundle, and a compliant housing. The cTAL’s fiber bundle has a resistance of approximately 0.3 mmHg/(L/min), making the combined resistance of the inlet and outlet sections around 0.2 mmHg/(L/min).

The compliant housing of the cTAL served to further lower resistance vs. identical hard shell designs (Figure 3.10).
The housing of the cTAL prototype provided a large compliance of $5.21 \pm 0.57 \text{ mL/mmHg}$ under high pressure ($> 5 \text{ mmHg}$). This compliance allowed the housing to expand, accommodating a larger volume of blood during systole and dampening the flow pulse. As the housing expands, fluid enters a larger area and the velocity of the fluid decreases. Minor losses are reduced and less recirculation occurs, lowering $Z_0$.

Comparison of cTAL vector plots with those in the hard shell housing TAL described in Chapter 2 confirmed that recirculation regions are smaller and have lower velocity in the cTAL. This resulted in lower impedances compared to the rigid housing TAL. At 4 L/min and 100 beats/min, $Z_0$ was $0.53 \pm 0.06 \text{ mmHg/(L/min)}$ and $Z_I$ was $0.86 \pm 0.08 \text{ mmHg/(L/min)}$ in the cTAL with $\theta = 45^\circ$. In the same device with a rigid housing,
$Z_0 = 0.69$ and $Z_1 = 1.22$ mmHg/(L/min) [5]. Moreover, unlike TALs with rigid housings, $Z_0$ decreased slightly with increasing $SV$ in the cTAL (Figure 3.10). At $SV = 0.075$, $Z_0$ for the cTAL and TAL were 0.41 ± 0.14 and 0.94 ± 0.09 mmHg/(L/min), respectively. Thus, at higher flow rates, such as those experienced during exercise, there is a larger advantage to using the cTAL.

The lower impedance of the cTAL has implications for use in vivo in both PA-PA and PA-LA attachment modes. PA-LA attachment would be advantageous in patients with pulmonary hypertension. Since the device has an impedance much lower than the natural lungs, blood would flow preferentially through the cTAL. This would allow significant unloading of the right ventricle and return PA pressures to normal. One can predict how much the cTAL would lower PA pressures by using the simple formula:

$$P_{PA} = CO \times R + P_{LA},$$

where $P_{PA}$ is the PA pressure, $CO$ is cardiac output, $R$ is the resistance, and $P_{LA}$ is the left atrial pressure. Assuming $PVR = 4$ mmHg/(L/min), $P_{LA} = 6$ mmHg, and a cTAL resistance of 1.4 mmHg/(L/min) for the TAL plus anastomoses (adding 0.87 mmHg/(L/min) for the anastomoses) [6], the resulting system resistance with the cTAL attached in parallel would be 1.0 mmHg/(L/min). Assuming $CO = 6$ L/min, $P_{PA}$ decreases from 30 mmHg prior to cTAL attachment to 12 mmHg. If the original PVR is 8 mmHg/(L/min), $P_{PA}$ is reduced from 54 mmHg to 13 mmHg.

If the cTAL were attached in series with the native lungs (PA-PA), it would likely result in similar PA pressure but a reduction in CO due to additive impedances and the increased work load of the RV. Relationships developed by Kim et al, Kuo et al, and Akay et al predict the percentage decrease in CO as a result of increases in $Z_0$ and $Z_1$.
[3,6,7]. The increase in pulmonary system impedance ($\Delta Z$) due to the cTAL impedance ($Z_{cTAL}$) can be calculated with the equation:

$$\Delta Z = Z_{cTAL} f$$  \hspace{1cm} (6)

where $f$ is the fraction of flow to the cTAL. If we assume $CO=6$ L/min and allow 4 L/min to travel through the cTAL then $f=2/3$. We will use the cTAL *in vitro* data at $Q=4$ L/min and $HR=100$ beats/min, and add either 1, 2, or 3 mmHg/(min) for the anastomoses resistance. Although Akay *et al* [6] calculated anastomoses’ resistance to be 0.87 mmHg/(L/min), this value can vary greatly from case to case and thus we will assume a possible range of values for our calculations. The Kim, Kuo, and Akay predictions for decrease in CO are shown in Table 3.3.

Table 3.3: Kim, Kuo, Akay and average predictions for percentage decrease in CO for three different anastomoses’ resistances for $f=2/3$.

<table>
<thead>
<tr>
<th>Anastomoses’ Resistance, mmHg/(L/min)</th>
<th>Kim</th>
<th>Kuo</th>
<th>Akay</th>
<th>Average</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>4.1</td>
<td>12.8</td>
<td>7.6</td>
<td>8.2</td>
</tr>
<tr>
<td>2</td>
<td>8.5</td>
<td>15.3</td>
<td>12.6</td>
<td>12.1</td>
</tr>
<tr>
<td>3</td>
<td>12.7</td>
<td>17.7</td>
<td>17.5</td>
<td>16.0</td>
</tr>
</tbody>
</table>

If the Kim and Akay data is averaged and we assume an anastomoses’ resistance of 1 mmHg/(L/min), $f=0.94$ results in a 10% decrease in CO. However, if we assume a higher anastomoses’ resistance of 3 mmHg/(L/min), in order to only get a 10% decrease in CO, $f$
must be \( \leq 0.46 \). Overall, these predictions are lower than previous devices and may allow for cTAL attachment in series in a small group of patients without significant pulmonary hypertension.

Gas transfer in the cTAL is excellent with outlet PO\(_2\) and SO\(_2\) values above 240 mmHg and 99.5\%, respectively, at each tested flow rate. Thus, the cTAL exceeds its gas transfer goals and the fiber bundle size could be reduced in the future. This could result in a smaller device, offering the advantage of a smaller prime volume and a lighter, more “wearable” device. A shortened device would also help to eliminate any low flow or stasis regions near the distal end.

In conclusion, this cTAL features excellent gas transfer and far lower impedance than any previous TAL. Its performance in vivo will be described in the following two chapters. PA-LA and PA-PA in vivo testing will examine the cTAL’s interaction with the right ventricle as well as its biocompatibility.
References


Chapter 4
In-Parallel cTAL Attachment

Introduction

The most commonly proposed TAL attachment modes are in parallel or in series with the natural lungs. During in parallel attachment blood flow is routed from the pulmonary artery (PA), through the TAL and then returned to the left atrium (LA). In patients with pulmonary hypertension this attachment mode is ideal, as it reduces $Z_0$ for the combined TAL and natural lung system and thus unloads the right ventricle. In this setting, TAL attachment should ideally result in a pulmonary system $Z_0$ that is as close as possible to the healthy natural lung. To accomplish this, the combined resistance of the TAL and the anastomoses used to attach it must also be close to that of the healthy natural lung.

The cTAL described in previous chapters was designed to meet this goal. This study examined pulmonary hemodynamics and cTAL function during in parallel attachment in healthy sheep. Device flows of 60-90% of CO through the cTAL were tested both at rest and during simulated exercise.
Methods

Compliant Thoracic Artificial Lung

The cTAL developed and described in Chapter 3 was used for in vivo testing (Figure 3.1). A holder was designed out of PVC to maintain the cTAL in an upright position, prevent the inlet and outlet conduits from kinking, and clamp the sides of the device to prevent blood from shunting around the fiber bundle (Figure 4.1).

![Figure 4.1: cTAL in holder for in vivo studies](image)

Experimental Procedure

A total of five male sheep averaging 60.2 ± 1.9 kg were used in this study. All sheep received humane care in compliance with the “Guide for the Care and Use of Laboratory Animals” and all methods were approved by the University of Michigan Committee for the Use and Care of Animals. Anesthesia was induced with 6-9 mL/kg of propofol and then maintained after endotracheal intubation with 1-3% inhaled isoflurane.
Sheep were mechanical ventilated with oxygen using a Narkomed 6000 ventilator (North American Dräger, Telford, PA). The ventilator was set to a tidal volume of 10 mL/kg and a frequency of 12 breaths/min and was adjusted to maintain an arterial partial pressure of carbon dioxide (PaCO₂) between 35 and 45 mmHg. A carotid arterial line and left jugular venous line were placed and then connected to fluid coupled pressure transducers (Hospira, Inc., Lake Forest, IL) for the continuous monitoring of arterial and central venous pressures (\(P_{\text{Art}}\) and \(P_{\text{CV}}\)).

A muscle sparing left thoracotomy was performed and the pulmonary artery and left atrium (LA) were identified. Dacron vascular grafts (Terumo, Ann Arbor, MI) for artificial lung attachment were attached to the PA and LA. An ultrasonic perivascular flow probe (Transonic 24AX, Transonic Systems, Inc., Ithaca, NY) was placed around the PA, proximal to the device inlet graft. This probe was connected to a flow meter (T400, Transonic Systems, Ithaca, NY) to allow for the measurement of PA flow (\(Q_{\text{PA}}\)). A pressure catheter (Becton Dickinson and Co., Franklin Lakes, NJ) was then inserted at the proximal PA and connected to a transducer for the display and recording of the PA pressure (\(P_{\text{PA}}\)). A Rommel tourniquet was placed around the distal PA to allow for the adjustment of flow through the cTAL. Prior to device attachment, 1g of methylprednisolone (Solu-Medrol, Pfizer, New York, NY) was administered and the animal was anticoagulated with 100 IU/kg of intravenous sodium heparin (Baxter Healthcare Corp., Deefield, IL) to maintain active clotting times of above 300 seconds.

The cTAL was primed with heparanized saline (10 U/mL) and then connected to the PA (device inlet) and LA (device outlet) grafts (Figure 4.2).
An ultrasonic flow probe (Transonic 14PXL, Transonic Systems, Inc., Ithaca, NY) was placed around the inflow conduit and connected to a flow meter (T400, Transonic Systems Inc., Ithaca, NY) in order to measure device flow ($Q_{cTAL}$). Pressure transducers were connected to the device inlet and outlet to acquire device inlet and outlet pressure ($P_{in}$ and $P_{out}$). A suction line was attached to the cTAL gas outlet and 95:5% O$_2$:CO$_2$ with 2-3% vaporized isoflurane between 3 and 9 L/min was used as the sweep gas through the gas inlet. Sweep gas flow was adjusted during cTAL use to maintain PaCO$_2$ between 35 and 45 mmHg. A Hoffmann clamp was placed around the cTAL outlet conduit to restrict flow through the device. After cTAL attachment, clamps on the cTAL inlet and outlet conduits were removed and the Hoffmann clamp was slowly loosened until $Q_{cTAL} = 1$ L/min was achieved. This flow was maintained for 10 minutes for equilibration of fluid volumes and any inflammatory response. After device attachment, CVP decreased related
to the systemic inflammatory response and 500 mL of hetastarch and 500 – 1000 mL of
crystalloid were administered to restore CVP back to baseline values. Thereafter, the
device conduits were clamped off and baseline data was taken, marking the start of the
experiment.

A hemodynamic data set of $Q_{PA}$, $Q_{cTAL}$, $P_{art}$, $P_{CV}$, $P_{PA}$, $P_{in}$, $P_{out}$ were digitally
acquired for 10 seconds at a sampling frequency of 250 Hz through a BIOPAC data
acquisition system (BIOPAC, Goleta, CA). Before attaching the cTAL, an animal
baseline data set was acquired. Rest and exercise conditions were simulated using a
continuous dobutamine (Hospira Inc., Lake Forest, IL) infusion of 0 and then 5
mcg/kg/min, respectively. At each dose, the hemodynamic data set was acquired at
baseline ($Q_{cTAL} = 0$ L/min) and conditions of 60, 75, and 90% of CO shunted to the cTAL
($100*Q_{cTAL}/Q_{PA}$), created by tightening the Rommel tourniquet around the PA. At each
condition, 10 minutes were allowed for equilibration before data was taken. Also, at each
condition, a device exit gas sample was taken along with blood samples from the animal,
device inlet and device outlet.

**Data Analysis**

The zeroth harmonic pulmonary input impedance modulus, $Z_0$, was calculated at
each flow condition:

$$Z_0 = \frac{P_0}{Q_0} \quad (1)$$

where $P_0$ is the mean PA pressure and $Q_0$ is the mean PA blood flow rate.

Device resistance, $R$, was calculated using the formula:
where $P_{in}$ is the cTAL inlet pressure, $P_{out}$ is the cTAL outlet pressure, and $Q_{cTAL}$ is cTAL flow. Data collected from device inlet and outlet blood gas samples were used to calculate oxygen transfer rates, VO$_2$, as done in Chapter 3 (equation 4). Comparisons were performed on the data, separately for rest and exercise sets, with SPSS 19 (SPSS, Chicago, IL). A mixed model was used with sheep number as the subject variable and flow condition (percentage of flow to the cTAL) as the fixed, repeated-measure variable. All data is reported as mean ± standard error with a p-value of 0.05 or less being considered statistically significant.

Results

Animal Physiology

Before cTAL attachment, average animal baseline CO, mean arterial pressure (MAP), mean PA pressure (mPAP), central venous pressure (CVP), and $Z_0$ was $6.4 ± 0.59$ L/min, $87.7 ± 8.08$ mmHg, $18.2 ± 1.34$ mmHg, $6.83 ± 1.22$ mmHg and $2.99 ± 0.41$ mmHg/(L/min), respectively. The effect of cTAL attachment on the sheep’s CO is shown in Figure 4.3.
Without dobutamine, the baseline CO was 5.5 ± 0.65 L/min and was maintained at or above this level until 90% flow to the cTAL. At 90% flow to the cTAL, there was a negligible, 0.33% decrease in CO from baseline which was not significant (p=0.94). At 5 mcg/kg/min of dobutamine, baseline CO was 7.23 ± 0.70 L/min and was maintained above this level until it decreased to 6.88 ± 0.69 L/min at 90% flow to the cTAL. This 5.6% drop was also not significant (p=0.36).

Figure 4.4 displays $Z_0$ for increasing flow to the cTAL.
For both rest and dobutamine-simulated exercise conditions, $Z_0$ decreases from baseline with 60% flow to the cTAL. Without dobutamine, $Z_0$ remains below baseline at all conditions. At 5 mcg/kg/min of dobutamine, baseline $Z_0$ decreases due to increased blood flow to the natural lungs. As more flow is diverted to the TAL, $Z_0$ is initially relatively constant but begins to rise at 75% flow. As the natural lung is increasingly excluded, $Z_0$ then rises towards the same $Z_0$ as without dobutamine. At 90% flow, $Z_0$ has increased 50% above the 5 mcg/kg/min dobutamine baseline and is almost equivalent to all 0 mcg/kg/min conditions. This increase was significant ($p=0.049$) when compared to baseline.
MAP and mPAP for both dobutamine levels are displayed in Table 4.1 at each target and actual percent flow to the cTAL.

Table 4.1: Mean pulmonary artery pressure and mean arterial pressure for varied percentages of the cardiac output to the cTAL

<table>
<thead>
<tr>
<th>Percentage of CO to cTAL</th>
<th>Dobutamine Dose</th>
<th>Goal</th>
<th>Actual</th>
<th>mPAP (mmHg)</th>
<th>MAP (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BL – 0%</td>
<td>0 mcg/kg/min</td>
<td>0 ± 0%</td>
<td>25.1 ± 5.3</td>
<td>78.4 ± 12.6</td>
<td></td>
</tr>
<tr>
<td>60%</td>
<td>61 ± 1%</td>
<td>21.6 ± 4.2</td>
<td>72.3 ± 6.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>75%</td>
<td>74 ± 2%</td>
<td>22.0 ± 4.6</td>
<td>73.5 ± 7.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>90%</td>
<td>91 ± 2%</td>
<td>24.3 ± 5.2</td>
<td>73.1 ± 7.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BL – 0%</td>
<td>5 mcg/kg/min</td>
<td>0 ± 0%</td>
<td>20.5 ± 1.5</td>
<td>78.5 ± 5.0</td>
<td></td>
</tr>
<tr>
<td>60%</td>
<td>61 ± 2%</td>
<td>20.5 ± 2.6</td>
<td>73.2 ± 2.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>75%</td>
<td>76 ± 2%</td>
<td>26.5 ± 3.2</td>
<td>73.7 ± 6.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>90%</td>
<td>93 ± 2%</td>
<td>29.4 ± 3.6</td>
<td>67.3 ± 6.9</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

BL = baseline; CO = cardiac output; cTAL = compliant thoracic artificial lung; MAP = mean arterial pressure; mPAP = mean pulmonary artery pressure

The mean baseline PA pressure at 0 mcg/kg/min dobutamine was 25.1 ± 5.31 mmHg with no significant change at 60, 75 and 90% flow to the cTAL (p = 0.35, 0.46, and 0.86). At 5 mcg/kg/min dobutamine, mPAP increased from 20.5 ± 1.48 mmHg at baseline to
29.4 ± 3.61 mmHg at 90% flow to the cTAL; however, this increase approached but was not significant (p=0.09). Baseline MAP was 78.4 ± 12.6 mmHg with no dobutamine and decreased slightly to 73.1 ± 7.82 mmHg at 90% flow to the cTAL. With 5 mcg/kg/min of dobutamine, baseline MAP was 78.5 ± 4.98 mmHg and decreased to 67.3 ± 6.92 mmHg at 90% flow to the cTAL; however, this decrease was not significant (p=0.16).

**Device Performance**

Device resistance at various flow rate ranges is shown in Figure 4.5.

![Figure 4.5: cTAL resistance at varying blood flow rate ranges](image)

Resistance of the cTAL remains relatively constant at all tested flow rates. Device resistance averaged 0.51 ± 0.01 mmHg/(L/min), ranging from 0.50 ± 0.01 mmHg/(L/min) at 2-3 L/min to a maximum of 0.55 ± 0.02 mmHg/(L/min) at 5-6 L/min. Accordingly,
device flow did not significantly effect resistance (p=0.29). Though this was not primarily a gas exchange experiment, the cTAL also exchanged gas effectively. Arterial PO$_2$, PCO$_2$ and pH, along with cTAL O$_2$ and CO$_2$ gas transfer rates (VO$_2$ and VCO$_2$) are displayed in table 4.2 for each device flow condition.

<table>
<thead>
<tr>
<th>Dobutamine Dose (mcg/kg/min)</th>
<th>% CO to cTAL</th>
<th>Arterial PO$_2$ (mmHg)</th>
<th>Arterial PCO$_2$ (mmHg)</th>
<th>Arterial pH</th>
<th>cTAL VO$_2$ (mL/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>60%</td>
<td>439 ± 57</td>
<td>38.9 ± 1.3</td>
<td>7.40 ± 0.01</td>
<td>85.7 ± 7.7</td>
</tr>
<tr>
<td>75%</td>
<td>460 ± 49</td>
<td>38.7 ± 1.3</td>
<td>7.41 ± 0.01</td>
<td>96.0 ± 6.8</td>
<td></td>
</tr>
<tr>
<td>90%</td>
<td>442 ± 35</td>
<td>37.1 ± 3.6</td>
<td>7.40 ± 0.01</td>
<td>109.1 ± 8.2</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>60%</td>
<td>399 ± 37</td>
<td>43.6 ± 1.6</td>
<td>7.37 ± 0.02</td>
<td>103.3 ± 11.1</td>
</tr>
<tr>
<td>75%</td>
<td>442 ± 35</td>
<td>45 ± 1.8</td>
<td>7.36 ± 0.02</td>
<td>120.7 ± 10.9</td>
<td></td>
</tr>
<tr>
<td>90%</td>
<td>422 ± 38</td>
<td>44.4 ± 1.3</td>
<td>7.36 ± 0.01</td>
<td>155.5 ± 8.8</td>
<td></td>
</tr>
</tbody>
</table>

CO = cardiac output; cTAL = compliant thoracic artificial lung; VO$_2$ = O$_2$ gas transfer rate
The device outlet oxyhemoglobin saturations were above 99% at every flow condition, although venous conditions were not sufficient to challenge the device. A typical hemoglobin and device inlet PO$_2$ was 7.5 and 80 mmHg, respectively. Even with this higher inlet PO$_2$, VO$_2$ rates ranged from $85.7 \pm 7.7$ mL/min at 0 dobutamine and 60% CO to the cTAL to $155.5 \pm 8.8$ mL/min at 5 dobutamine and 90% CO to the cTAL. Lastly, at the conclusion of the experiment, there was no visible clot formation in the device.

Discussion

To date, there is no commercially available TAL. Over 18 years of development, these devices have always featured excellent gas exchange and research has thus focused on developing devices with progressively lower blood flow resistance and improved hemodynamics during *in vivo* attachment [Cook, Cook, Sato]. Ideally, TALs should be able to provide the majority of the gas exchange while being able to maintain normal pulmonary hemodynamics.

As there is no commercial device for this application, a few groups have investigated clinical PA-LA attachment in cases of severe pulmonary hypertension using the lowest resistance gas exchanger on the market, the Novalung ILA. Two of these patients had primary pulmonary hypertension and four had pulmonary veno-occlusive disease. Five of the patients (87%) were successfully bridged [Schmid, Camboni, Strueber]. Despite this success, the Novalung was not designed for this application. Its intended use is arterio-venous CO$_2$ removal, which features much lower blood flows of 2 L/min and higher driving pressures. As a result, the Novalung’s gas exchange capabilities
are low for the application and the resistance is approximately 5-6 mmHg/(L/min) at blood flows of 2-2.5 L/min [Florchinger, Muller, Wiebe]. To support COPD, IPF, and septic patients, greater gas exchange will be required. Moreover, in cases of marked pulmonary hypertension, it would be ideal to be able to fully unload the RV and eliminate the need for inotropes.

In the current study, the new cTAL design was tested in healthy animals with normal PA pressure. Therefore, the goal for this study was to maintain normal PA pressures and CO with up to 90% of CO diverted to the cTAL. Results indicate that these goals were met. cTAL attachment caused only minimal, statistically insignificant decreases in CO and increases in PA pressure from baseline at all conditions. At the most extreme condition, simulated exercise with 90% of the CO diverted to the cTAL, CO decreased 5.6% and PA pressure increased 8.8 mmHg. This, however, represents an extreme case. A more typical, and advisable [Takewa], condition would likely be 75% of CO. Here, CO was identical to baseline values, and PA pressure decreased at rest and increased a small amount during simulated exercise.

Maintenance of normal pulmonary hemodynamics was the result of very low cTAL resistance. Average resistance for the range of flows tested was 0.51 ± 0.01 mmHg/(L/min), well below that of the natural lung. Moreover, the resistance of the device did not change significantly as flow increased. The low cTAL resistance, in turn, maintains low $Z_0$. Previous studies have shown that $Z_0$ is the dominant variable affecting CO during TAL attachment, with CO decreasing as $Z_0$ increases [Kim, Kuo, Akay]. In this study, $Z_0$ is lower than baseline at 60% flow to the cTAL due to the second, parallel flow path provided by the cTAL and the minimal banding of the PA at that condition. As
the PA is banded further, the natural lung portion of the system is closed, the percentage of flow to the cTAL increases, and \( Z_0 \) increases slightly. However, since device resistance remains low, the resulting \( Z_0 \) remains small and similar to that of healthy natural lungs. At 5 mcg/kg/min of dobutamine, however, high CO and PA pressure leads to lower baseline \( Z_0 \). As flow is diverted to the cTAL, \( Z_0 \) then increases to the same level seen with in the 0 dobutamine case.

Based on these results, the cTAL will be able to completely unload the right ventricle in vivo during PA-LA attachment in patients with any degree of pulmonary hypertension. To examine this, one can use the equation \( P_{PA} = CO \cdot R + P_{LA} \), where \( P_{PA} \) is the PA pressure, \( P_{LA} \) is the left atrial pressure, and \( CO \) is cardiac output. \( R \) is the resistance of the parallel artificial and natural lung system. If there is no PA banding, \( R = [R_N \cdot (R_T + R_A)]/[R_N + R_T + R_A] \), in which \( R_N \) is the natural lung resistance, \( R_T \) is the TAL resistance, and \( R_A \) is the resistance of the TAL anastomoses or cannulae. For \( CO = 6 \) L/min, \( P_{LA} = 6 \) mmHg, \( R_T = 0.5 \) mmHg/(L/min), and \( R_A = 0.87 \) mmHg/(L/min), and \( R_N = 9 \) mmHg/(L/min) [10], the PA pressure prior to attaching the TAL would be 60 mmHg. With the cTAL, it falls to 13 mmHg. In comparison, under the same conditions, the Novalung ILA (\( R_T = 6 \) mmHg/(L/min)) would reduce PA pressure to 29 mmHg.

The other experimental TAL in development is the Biolung® (MC3, Ann Arbor, MI). The hard-shell Biolung was tested in a study similar to this one [Akay 2]. The only exception is that Akay et al simulated rest (no dobutamine), ambulatory (2 mcg/kg/min dobutamine), and exercise (5 mcg/kg/min dobutamine) conditions. Results showed that CO was maintained as the percentage of CO to the device increased at resting and ambulatory conditions. At exercise conditions, CO decreased with increasing flow to the
TAL up to 23 ± 5% at 90% flow diverted through the TAL. Direct comparison between these two studies is difficult, as the baseline $Z_0$ and baseline cardiac response to dobutamine varied greatly. In the Biolung study, baseline $Z_0$ were lower and, as a result, baseline CO was greater at the same dobutamine doses. Thus, the easiest means of comparison between these studies is the $Z_0$ with 90% of cardiac output through the artificial lung. Here, hemodynamics are largely unaffected by natural lung resistance. At 90%, $Z_0$ ranged from 4.5-4.9 in the Biolung study and 4.1-4.5 in the cTAL study. This small difference is similar to what one would predicted by the difference in TAL resistances in these studies. The cTAL resistance is approximately 0.3 mmHg/(L/min) smaller with flow rates of 2-3 L/min and approximately 0.65 mmHg/(L/min) lower at flow rates of 5-6 L/min. Thus the Biolung should provide slightly less but similar unloading.

Ultimately, this study suggests that the cTAL is capable of being used clinically, in parallel with the native lungs and under high flow conditions. Minimal decreases in CO were seen at 90% flow to the device at both dobutamine levels, indicating exercise would be possible during cTAL attachment. Since cTAL resistance is small, flow should route preferentially through the device. This attachment mode would be ideal for patients with high pulmonary vascular resistance since the device resistance would be much lower than the native lung resistance. Approximately 57% of CO went through the cTAL with no PA banding during this experiment. Assuming a pulmonary vascular resistance of 2.25 mmHg/(L/min) in patients with pulmonary hypertension, about 70% of the CO would flow preferentially through the cTAL with no PA banding. Further testing of in parallel attachment of the cTAL is required in sheep with pulmonary hypertension. The long term
effects of in parallel cTAL attachment will also be studied for up to 30 days in healthy sheep and sheep with pulmonary hypertension.

**Conclusion**

The cTAL resistance is lower than the natural lung, averaging $0.51 \pm 0.03$ mmHg/(L/min). Use of the cTAL in a PA-LA configuration caused no change in CO under rest and exercise conditions. Thus, the cTAL can provide PA-LA respiratory support without significant changes in pulmonary hemodynamics in healthy sheep and will be capable of unloading the RV in subjects with pulmonary hypertension.
References


Chapter 5

In-Series cTAL Attachment

Introduction

While the cTAL had excellent *in vivo* performance in parallel, there is the constant risk of life-threatening air emboli and thromboemboli in this attachment mode. An alternative configuration is in series attachment with the natural lungs. In this mode, blood is routed from the proximal pulmonary artery (PA) through the cTAL and then back to the distal PA where the blood then flows through the native lungs and finally back to the LA. This configuration eliminates the risk of systemic emboli by allowing the lung to perform its filtration and non-respiratory functions. In addition, the entire cardiac output (CO) can flow through the cTAL, allowing for complete gas exchange. However, studies on TAL attachment mode have shown series attachment to increase pulmonary system impedance and decrease CO [1]. Increased pulmonary system impedance is caused by the additive impedance of the TAL and natural lung. Therefore, it is imperative that the device have the lowest impedance possible to prevent RV failure when attached in series.

The low-resistance cTAL developed and tested in previous chapters was examined in this setting. This study examined pulmonary hemodynamics and cTAL function during in series attachment in healthy sheep. As in the PA-LA study, described
in Chapter 4, different device flows were tested with 60-90% of CO through the cTAL both at rest and during simulated exercise.

Methods

Experimental Procedure

Five male sheep averaging 63 ± 0.92 kg were used in this study. All sheep received humane care in compliance with the “Guide for the Care and Use of Laboratory Animals” and all methods were approved by the University of Michigan Committee for the Use and Care of Animals. Anesthesia was induced with 6-9 mL/kg of propofol and then maintained after endotracheal intubation with 1-3% inhaled isoflurane (Abbot Laboratories, Chicago, IL). Sheep were mechanically ventilated with oxygen using a Narkomed 6000 ventilator (North American Dräger, Telford, PA). The ventilator was set to a tidal volume of 10 mL/kg and a frequency of 12 breaths/min and was adjusted to maintain an arterial PCO2 (PaCO2) between 35 and 45 mmHg. A carotid arterial line and left jugular venous line were placed and then connected to fluid coupled pressure transducers (ICU Medical, Inc., San Clemente, CA) for the continuous monitoring of arterial and central venous pressures (P_{Art} and P_{CV}).

A muscle sparing left anterolateral thoracotomy was performed and the fourth rib was removed. The left lung was packed laterally and the pericardium was incised. The PA was then identified. Low porosity woven Dacron vascular grafts 18 mm in size (Boston Scientific, Natick, MA) solvent bonded to 5/8” inner diameter PVC tubing (Fisher Scientific, Pittsburgh, PA) were used as conduits for artificial lung attachment. An end-to-side anastomosis was performed between the device outlet conduit and the
distal PA near the bifurcation. The device inlet end-to-side anastomosis was then performed on the proximal PA. An ultrasonic perivascular flow probe (Transonic 24AX, Transonic Systems, Inc., Ithaca, NY) was placed around the PA, proximal to the device inlet graft. This probe was connected to a flow meter (T400, Transonic Systems, Ithaca, NY) to allow for the measurement of PA flow ($Q_{PA}$). A pressure catheter (Becton Dickinson and Co., Franklin Lakes, NJ) was then inserted at the proximal PA and connected to a transducer for the display and recording of the proximal PA pressure ($P_{PAP}$). A Rommel tourniquet was placed around the PA between the inlet and outlet grafts to allow for the adjustment of flow through the cTAL. Pressure catheters were inserted into the distal PA near the bifurcation and into the left atrium to allow for the recording of the distal PA pressure ($P_{PAD}$) and the left atrial pressure ($P_{LA}$), which facilitated the calculation of the pulmonary vascular resistance (PVR) of the native lungs. Prior to device attachment, 1g of methylprednisolone (Solu-Medrol, Pfizer, New York, NY) was administered, and the animal was anticoagulated with 100 IU/kg of intravenous sodium heparin (Baxter Healthcare Corp., Deerfield, IL) to maintain active clotting times of above 300 seconds.

The cTAL was primed with heparinized saline (10 U/mL) and then connected to the proximal PA (device inlet) and distal PA (device outlet) grafts (Figure 5.1).
An ultrasonic flow probe (Transonic 14PXL, Transonic Systems, Inc., Ithaca, NY) was placed around the inflow conduit and connected to a flow meter (T400, Transonic Systems Inc., Ithaca, NY) in order to measure device flow \(Q_{cTAL}\). Device inlet and outlet pressure \((P_{in} \text{ and } P_{out})\) were measured by fluid-coupled pressure transducers. A suction line was attached to the cTAL gas outlet and 95:5% O\(_2\):CO\(_2\) with vaporized isoflurane (1-3%) was used as the sweep gas through the gas inlet. Sweep gas flow was adjusted during cTAL use to maintain PaCO\(_2\) between 35 and 45 mmHg. A Hoffmann clamp was placed around the cTAL outlet conduit to restrict flow through the device. After cTAL attachment, clamps on the cTAL inlet and outlet conduits were removed and the Hoffmann clamp was loosened until 1 L/min flow to the cTAL was achieved. This flow was maintained for 10 minutes for equilibration of fluid volumes and any inflammatory response. After device attachment, CVP decreased related to the systemic inflammatory response and up to 500 mL of hetastarch and 500 mL of crystalloid were administered to

Figure 5.1: PA-PA cTAL attachment and instrumentation
restore CVP back to baseline values, if required. Thereafter, the device conduits were clamped off and baseline data was taken.

A hemodynamic data set of $Q_{PA}$, $Q_{cTAL}$, $P_{art}$, $P_{CV}$, $P_{PA}$, $P_{in}$, $P_{out}$ were digitally acquired for 10 seconds at a sampling frequency of 250 Hz through a BIOPAC data acquisition system (BIOPAC, Goleta, CA). Rest and exercise conditions were simulated using a continuous dobutamine (Hospira Inc., Lake Forest, IL) infusion of 0 and then 5 mcg/kg/min, respectively. At each dose, the hemodynamic data set was acquired at baseline, with no flow to the device, and conditions of 60, 75, and 90% of CO shunted to the cTAL ($Q_{cTAL}/Q_{PA}$), created by tightening the Rommel tourniquet around the PA. At each condition, 10 minutes were allowed for equilibration before data was taken. Also, at each condition, a device exit gas sample was taken along with blood samples from the animal, device inlet and device outlet to analyze the gas exchange properties of the device.

**Data Analysis**

The zeroth harmonic pulmonary input impedance modulus, $Z_0$, was calculated at each flow condition (equation 1, Chapter 4). $Z_0$ has been shown previously by our group to be an excellent index for predicting right ventricular function in high afterload states [2,3]. Device resistance, $R_{cTAL}$, was also calculated at each flow condition (equation 2, Chapter 4). The PVR of the native lungs was calculated by the following formula:

$$PVR = \frac{P_{PAD} - P_{LA}}{Q_0}$$  \hspace{1cm} (1)
where $P_{PAP}$ is the mean distal PA pressure, $P_{LA}$ is the mean left atrial pressure, and $Q_0$ is the mean PA blood flow rate (CO).

The resistance of the graft anastomoses, $R_a$, was calculated by analyzing all the resistive elements in the experimental pulmonary system setup (Figure 5.2).

![Figure 5.2: Experimental pulmonary system setup with all resistive elements. The shunt includes the anastomoses ($A_{in}$ and $A_{out}$), the cTAL and the flow occluder.](image)

Total pulmonary system resistance, $R_T$, was defined as:

$$R_T = \frac{P_{PAP} - P_{LA}}{Q_0} \tag{2}$$

where $P_{PAP}$ is the mean proximal PA pressure. Shunt resistance, $R_s$, was calculated by assuming it was an element in series with the native lungs:

$$R_s = R_T - PVR \tag{3}$$

By assuming the anastomoses were a resistive element in series with the cTAL, which was in parallel with the flow occluder, the following relationship was determined:
\[
\frac{1}{R_s} = \frac{1}{R_{cTAL} + R_a} + \frac{1}{R_{FO}} \tag{4}
\]

where \( R_{FO} \) is the resistance of the flow occluder defined using a flow relationship:

\[
R_{FO} = \frac{f(R_{cTAL} + R_a)}{1 - f} \tag{5}
\]

where \( f \) is the fraction of \( Q_0 \) diverted to the cTAL. Using equations 4 and 5, an equation for \( R_a \) was determined:

\[
R_a = \frac{R_s}{f} - R_{cTAL} \tag{6}
\]

\( R_a \) was calculated at each flow condition and an equation for approximate \( R_a \) as a function of \( f \) was determined.

Statistical analysis was performed separately for the 0 and 5 mcg/kg/min data sets. The analysis of baseline PVR before and after cTAL attachment was conducted using a paired two-tailed student’s t-test assuming equal variances. All other statistical comparisons were performed using a mixed model with sheep number as the subject variable and flow condition as the fixed, repeated-measure variable. The statistical analysis was performed using SPSS 19 (SPSS, Chicago, IL). All data is reported as the mean ± the standard error and a p-value of 0.05 or less was considered statistically significant.

**Results**

**Animal Physiology**

Before cTAL attachment, average animal baseline CO, mean arterial pressure (MAP), mean PA pressure (mPAP), central venous pressure (CVP), PVR, and \( Z_0 \) was 6.5
± 0.56 L/min, 87.6 ± 6.20 mmHg, 25.6 ± 0.90 mmHg, 12.1 ± 1.17 mmHg, 1.62 ± 0.32 mmHg/(L/min), and 4.05 ± 0.39 mmHg/(L/min), respectively. The effect of cTAL attachment on the sheep’s CO is shown in Figure 5.3.

Without dobutamine, the baseline CO of the sheep following cTAL attachment was 6.20 ± 0.53 L/min. This decreased 13% to 5.40 ± 0.34 L/min at 60% flow to the cTAL (p=0.922), 25% to 4.66 ± 0.31 L/min at 75% flow to the cTAL (p=0.07), and 35% to 4.05 ± 0.27 L/min at 90% flow to the cTAL (p=0.01). The CO did not drop significantly until greater than 75% of CO was diverted to the cTAL. After starting dobutamine at 5 mcg/kg/min, CO increased to 7.85 ± 0.70 L/min at baseline with no flow to the cTAL. Once again, CO decreased with increasing flows to the cTAL, dropping 4.9% to 7.46 ±
0.55 L/min, 12% to 6.93 ± 0.51 L/min, and 24% to 5.96 ± 0.44 L/min with 60, 75%, and 90% of CO going through the cTAL, respectively. These decreases in CO were not significant at any of the conditions (p=0.13).

Similarly, MAP decreased with increasing flow to the TAL (Figure 5.4).

![Figure 5.4: Mean arterial pressure, MAP, at varying percentages of cardiac output diverted to the cTAL for dobutamine doses of 0 and 5 mcg/kg/min.](image)

At 0 mcg/kg/min dobutamine, baseline MAP was 81 ± 4.0 mmHg and increased slightly to 83 ± 5.7 mmHg at 60% flow to the cTAL before decreasing to 76 ± 6.1 mmHg and 67 ± 5.0 mmHg at 75% and 90% flow to the cTAL, respectively (p=0.18). With 5 mcg/kg/min of dobutamine, baseline MAP was 84 ± 5.3 mmHg. This decreased to 80 ± 6.9 mmHg at 60% flow to the cTAL (p=0.99), 69 ± 2.8 mmHg at 75% flow to the cTAL.
(p=0.22), and then fell significantly to 60 ± 1.3 mmHg at 90% flow to the cTAL (p=0.01).

$Z_0$ increased with increasing flow to the cTAL (Figure 5.5).

![Figure 5.5: Zeroth harmonic impedance modulus, $Z_0$, at varying percentages of cardiac output diverted to the cTAL for dobutamine doses of 0 and 5 mcg/kg/min](image)

Without dobutamine, baseline $Z_0$ was 4.66 ± 0.56 mmHg/(L/min) and increased to 7.02 ± 0.56 mmHg/(L/min) at 75% flow to the cTAL (p=0.06) and 8.35 ± 0.67 mmHg/(L/min) at 90% flow to the cTAL (p<0.01). Baseline $Z_0$ in the simulated exercise condition was 4.27 ± 0.77 mmHg/(L/min) which slightly increased to 5.11 ± 0.19 mmHg/(L/min) at 75% flow to the cTAL (p=0.99). $Z_0$ increased significantly to 6.48 ± 0.45 mmHg/(L/min) once 90% flow to the cTAL was achieved (p<0.05).
Although $Z_0$ increased with increasing levels of PA occlusion, the mPAP (proximal) remained fairly stable under all of the rest and exercise conditions due to decreasing CO (Figure 5.6).

![Figure 5.6: Mean proximal pulmonary artery pressure, mPAP, at varying percentages of cardiac output diverted to the cTAL for dobutamine doses of 0 and 5 mcg/kg/min.](image)

Baseline mPAP without dobutamine was $28 \pm 2.0$ mmHg and increased by an insignificant amount to $33 \pm 2.5$ mmHg at 90% flow to the cTAL ($p=0.53$). Similarly, the baseline mPAP of $32 \pm 4.8$ mmHg increased insignificantly to $38 \pm 0.6$ mmHg at 90% flow to the cTAL under simulated exercise conditions ($p=0.57$).

The decrease in CO seen in Figure 5.3 is due to an increase in total pulmonary system resistance, $R_T$. An increase in $R_T$ at each condition is either due to an increase in PVR or everything prior to the native lungs (shunt resistance), $R_s$. The contribution of PVR and $R_s$ to $R_T$ is displayed in Figure 5.7.
Figure 5.7: Total pulmonary system resistance, $R_T$, pulmonary vascular resistance, PVR, and shunt resistance, $R_s$, at varying percentages of cardiac output diverted to the cTAL for dobutamine doses of (a) 0 mcg/kg/min and (b) 5 mcg/kg/min.
Baseline PVR, before cTAL attachment, was $1.61 \pm 0.32$ mmHg/(L/min). After attaching the cTAL, allowing flow to the device for 10 minutes, and then clamping off the device for baseline data, PVR increased substantially to $3.13 \pm 0.17$ mmHg/(L/min) ($p=0.01$). However, no further significant increase was seen with increasing percentage of CO to the cTAL ($p=0.10$). After the initiation of dobutamine, PVR decreased to $2.23 \pm 0.45$ mmHg/(L/min) at baseline. This represented a substantial drop from the baseline of the rest condition, although this did not achieve statistical significance ($p=0.16$). Again, PVR was unchanged as the percentage of CO to the cTAL was increased. However, as the percentage of CO to the cTAL increases, $R_s$ increases significantly at both rest and exercise conditions ($p<10^{-3}$). From Figure 5.7 it is apparent that the elevation in $R_T$ with increasing flow to the cTAL is due to $R_s$. Increases in $R_s$ must be due to either an increase in the resistance of the anastomoses, $R_a$, or the device resistance, $R_{cTAL}$ (Figure 5.8).
As percentage of CO to the cTAL increases, $R_a$ increases. A linear relationship between $f$ and $R_a$ was determined:

$$R_a = 1.77f + 1.31, \ R^2 = 61.7\%$$

$R_{cTAL}$ does not increase with increasing percentage of CO to the cTAL. Thus, increases in shunt resistance are due to increases in $R_a$ and not $R_{cTAL}$. Overall, $R_a$ is 4-6 times larger than the cTAL resistance and is thus the dominant cause of reduced cardiac output in the in-series configuration.

Figure 5.8: cTAL resistance, $R_{cTAL}$, and anastomoses’ resistance, $R_a$, at varying percentages of cardiac output diverted to the cTAL for dobutamine doses of 0 and 5 mcg/kg/min.
Device Function

The resistance of the cTAL was also assessed as a function of overall flow rate through the device (Figure 5.9). $R_{CTAL}$ did not change significantly with flow rate ($p=0.16$), averaging $0.46 \pm 0.02$ mmHg/(L/min) over the entire range of flows.

![Figure 5.9: cTAL resistance at varying blood flow rate ranges](image)

The cTAL also performed well in terms of gas transfer and biocompatibility. Due to relatively high venous saturations (approximately 85%) and lower hemoglobin ($6.11 \pm 0.17$ g/dL), the gas transfer function of the device was never challenged. Arterial blood gas hemoglobin oxygen saturations were maintained at near 100% for all of the conditions. Finally, all of the devices functioned without failure for the entire duration (3-4 hours) of the experiment and there was no visible clot formation within the device at the conclusion of the experiment.
Discussion

As discovered in Chapter 4, cTAL resistance did not change significantly with flow. In the current study, average cTAL resistance was 0.46 ± 0.02 mmHg/(L/min), similar to results from Chapter 4. Though cTAL resistance did not increase with increasing percentage flow to the device, $Z_0$ did increase. This increase was due to an increase in the shunt resistance and not PVR. There was an initial 94% increase in PVR from animal baseline during an initial equilibration phase following attachment of the cTAL due to an inflammatory response to the device. However, after this initial increase, the PVR did not increase significantly as percentage flow to the cTAL increased. Resistance of the shunt did, however, increase with increasing flow. Thus, increases in pulmonary system resistance with increasing flow diversion to the cTAL were due to increasing resistance of the anastomoses. These values were higher than predicted previously [4] but anastomoses’ resistance can vary greatly from case to case. Theoretically, at baseline (no flow to the cTAL), $R_T$ should have equaled PVR; however, this was not the case. There was a small additional resistance in the system, most likely from the anastomoses, suggesting the anastomoses cannot be treated as an element completely in series with the cTAL.

Based on previous predictions and the results in this chapter, we can make predictions on how much flow through the cTAL could be tolerated by patients. As discussed in earlier chapters, previous studies concluded that $Z_0$ is the dominant variable affecting CO during TAL attachment, with CO decreasing as $Z_0$ increases [2-4]. This was again found to be true in this study. For both dobutamine levels, as the percentage of CO
to the cTAL increased, $Z_0$ increased, causing CO to decrease. CO only decreased significantly at 90% of CO to the cTAL with 0 mcg/kg/min dobutamine. However, at the other conditions, there were percent decreases in CO that would not be acceptable for patient use. At 60% flow to the cTAL, with 0 and 5 mcg/kg/min, CO decreased 13% and 5%, respectively. Therefore, cTAL flows above 60% of CO are not advisable during PA-PA attachment in patients. From their study on the effects of impedance during PA-PA attachment, Akay et al determined that TAL resistance must be less than 1.81, 0.92, and 0.47 mmHg/(L/min) for 50, 75, and 100% of CO to the TAL to limit the drop in CO to less than 10%. However, they also calculated resistance of the anastomoses to be 0.87 mmHg/(L/min). In our study, the anastomoses resistance went up with increasing flow to the device and averaged 2.64 mmHg/(L/min). In Chapter 3, it was predicted that with an anastomoses’ resistance of 3 mmHg/(L/min), the percentage flow to the cTAL must be less than 46% for a decrease in CO less than 10%. Using the average cTAL and anastomoses’ resistance presented in this chapter, the percentage of CO to the cTAL must be \( \leq 53\% \) to maintain CO within 10% of baseline. Therefore, if anastomoses’ resistance is high, as in this study, the percentage of CO to the cTAL during PA-PA attachment should be kept around 50%. This warrants further examination in long-term cTAL studies. Results in these studies may vary due to additional variables, including anesthesia, animal health, and the length of cTAL attachment.
References


Chapter 6

Conclusion

Conclusions

This work examined the effect of different housing geometries and materials on TAL impedance and flow uniformity. From CFD and *in vitro* studies, the optimal TAL geometry was determined. A cTAL with this design was prototyped for *in vitro* and *in vivo* testing. The *in vivo* performance of the optimized cTAL was tested in two different attachment modes under simulated rest and exercise conditions. From these studies, it can be concluded that:

1. More gradual device entrances and exits result in reductions in TAL impedance. Recirculation in the inlet housing region is reduced as $\theta$ decreases, reducing overall device impedance. This was also the case for the cTAL, as confirmed by FSI analysis.

2. A region of little to no flow exists near the distal end of the TAL and cTAL in the 15° and 45° models. These areas were largest in the 15° model where flow was barely reaching past the middle of the housing body. This low flow region could lead to reduced gas transfer performance and clot formation over time.

3. Flow is uniform through the fiber bundle, with a resistance of about 0.30 mmHg/(L/min); approximately 40% of TAL resistance and 60% of cTAL resistance.
4. Average housing displacement was greatest in the 15° model. Thus, more flow dampening occurred for smaller $\theta$. The velocity entering the bundle at peak systole was highest in the 90° model, indicating that flow was less temporally uniform in this model due to lower dampening of the flow pulse. Low flow during diastole could also lead to stasis and clot formation.

5. The 45° model had the optimal combination of low impedance, better flow patterns, and relatively compact size. Thus, it was chosen for the cTAL prototype and tested *in vivo*.

6. The effect of flow rate on impedance was different in the hard-shell TAL and the cTAL. TAL $Z_0$ increased linearly with $SV$, while cTAL $Z_0$ decreased slightly with increasing $SV$. The cTAL had lower impedance than the TAL, especially at higher $SV$. Thus, at higher flow rates, such as those experienced during exercise, there is a larger advantage to using the cTAL.

7. Inlet flow pulsatility does not greatly influence TAL or cTAL impedance.

8. The cTAL has a large compliance ($5.21 \pm 0.57$ mL/mmHg under high pressure) which allows the housing to expand and dampen the flow pulse. Due to this expansion, the velocity of the fluid in the cTAL decreases, resulting in less recirculation and, as a result, lower $Z_0$. Comparison of cTAL vector plots with those in the hard shell housing TAL confirmed that recirculation regions are smaller and have lower velocity in the cTAL.

9. The cTAL has excellent gas transfer with outlet PO$_2$ and SO$_2$ values above 240 mmHg and 99.5%, respectively, at flow rates of 3-7 L/min. Thus, the rated flow of the device is well over 7 L/min.
10. *In vivo*, cTAL resistance does not vary significantly with blood flow rate. During PA-LA and PA-PA attachment, cTAL resistance averaged $0.51 \pm 0.03$ mmHg/(L/min) and $0.46 \pm 0.03$ mmHg/(L/min), respectively.

11. PA-LA cTAL attachment allows for excellent exercise tolerance and will be capable of unloading the RV in patients with pulmonary hypertension. Under all flow conditions during acute *in vivo* testing, CO was not significantly different from baseline. The cTAL should be able to provide PA-LA respiratory support for flows up to 90% of CO.

12. PA-PA attachment should be feasible at cTAL blood flows up to 53% of CO.

**Limitations and Future Work**

One limitation in this work is the ability of the CFD software to accurately model the experimental flow conditions. As discussed in Chapters 2 and 3, turbulent flow could not be applied in the fiber bundle and thus parts of the models had to be run separately. This worked well in the hard-shell TAL study, with CFD results matching experimental results fairly closely (Chapter 2). However, FSI analysis proved to be more complicated and a different approach was necessary to achieve physically accurate results. Because the outlet housing section had to be run as laminar, impedance values were higher overall in the FSI models, creating slight differences from the cTAL experimental results (Chapter 3). Despite these challenges, the CFD studies provided important information on the flow uniformity within the device throughout the cardiac cycle. Velocity plots revealed areas of little to no flow at the distal end of the device. Since the cTAL exceeded its gas transfer goals, the fiber bundle size could be reduced in the future. A shortened
device would help to eliminate any low flow or stasis regions near the distal end. A smaller device would also offer the advantages of a smaller prime volume and a lighter, more “wearable” device.

For the current cTAL design, the next step is 14 day *in vivo* studies in both healthy and pulmonary hypertension sheep models. Two long-term experiments have already been carried out in healthy sheep. In these studies, the cTAL was attached in the PA-LA configuration with no PA banding. After the animal recovered from surgery, the cTAL was placed in a modified backpack, attached to the sheep’s flank, and attached to the inlet and outlet grafts (Figure 6.1).

![Figure 6.1: cTAL and device holder inside a modified backpack, attached to the sheep’s flank.](image)

In the first sheep, the cTAL was attached for 16 days. The second sheep was euthanized after three days due to inlet graft occlusion from sheep positioning and movement. Gas
transfer of the device was excellent and cTAL resistance did not change significantly over the course of either experiment. In the first study, resistance started at 0.95 mmHg/(L/min) and averaged 0.66±0.04 mmHg/(L/min) over the entire 16 days. At the conclusion of this experiment, the device was examined for blood clots. No clots were present, except for the sides of the bundles where it was clamped in the device holder (Figure 6.2).

Figure 6.2: Clotting on the side of the fiber bundle where the cTAL was clamped in its device holder.

Clamping tightly on the sides of the bundle, to prevent shunting and keep the device firmly in place, created a static region where blood pooled. Also, the cTAL is positioned in the backpack such that it lies on its side, with one side of the bundle higher than the other. More clotting was present on the highest side due to less blood reaching that area. In the second sheep, modifications were made to the side clamps so they contacted less of the bundle. The sides were also clamped more lightly. Though attachment only lasted three days, this appeared to decrease clotting in the side regions.
In the future, the cTAL holder will be redesigned so the device is held in place via attachments to the gas caps. The minimum amount of clamping necessary to prevent shunting will be then be applied to the bundle sides. The possibility of creating internal means of preventing shunting, which would eliminate the need for any side clamping, is being investigated. Ultimately, long-term \textit{in vivo} testing will reveal additional information on device longevity and tolerance in different attachment configurations, as well as different disease models.

Figure 6.3: cTAL from 3 days of attachment in a chronic sheep, using more moderate clamping of the sides. No clotting in the cTAL body (a) and reduced clotting in the side regions (b)
Appendix

Compliant Thoracic Artificial Lung Fabrication

Summary

The cTAL models used for testing in Chapters 3-5 in this dissertation were constructed with a multi-step fabrication process. The compliant cTAL housing is created by dipping a 4-part Delrin model (Figures A.1 and A.2) in a container of Biospan. After this 6 day process, the housing is removed from the dipping model and goes through an extensive process of testing for holes/leaks and then patching the housing. cTAL fiber bundles are created by winding a mat of polypropylene hollow fibers around a polycarbonate core. The completed fiber bundle is then inserted into the center of the patched housing. After trimming excess housing, the cTAL is placed in our centrifuge setup for potting: binding the fiber bundle and housing together. A two part polyurethane polymer, WC7-53 (BJB Enterprises Inc., Tustin, CA), is injected through the top of the spinning centrifuge which evenly distributes the material into the ends of the housing/fiber bundle and binds them together. After potting, the ends of the cTAL are cut to expose the open ends of the hollow fibers. The cTAL then undergoes another series of leak testing and patching to eliminate any holes and/or leaks. Acrylic gas caps are then adhered to both ends of the fiber bundle with silicone sealant. Finally, the cTAL is tested
for air leaks. After these are patched, the cTAL is ready for testing. Detailed protocols are included below for each part of the cTAL fabrication process.

**Dipping: Creating the cTAL Housing**

**Percent Solid**

Before beginning the dipping process, the Biospan must be checked for the correct percent solid formulation as described below.

1. Weigh a weigh dish three times, zeroing after each time.
2. Average the three masses and record them on the percent solid sheet in the front of the dipping log.
3. Use a 3 ml syringe to collect 1.5 – 2.0 ml of BioSpan from the hood.
4. Dispense BioSpan into weigh dish and repeat steps 1 and 2.
5. Keep the BioSpan in the oven (at 120˚ F) overnight.
6. Repeat steps 1 and 2 with the cured BioSpan.
7. Subtract the weight of the weigh dish from the uncured BioSpan and cured BioSpan masses. Divide the weight of the cured BioSpan by the weight of the uncured BioSpan and multiply by 100. If the percent is 20 ± .5%, then congratulations, you don’t have to add anything! If the percent is outside the .5% range, then you either have to add DMAC or BioSpan.

*(Warning: whenever the vat of BioSpan is open or there is something being dipped, keep the attached hood closed and make sure the hose is attaching both hoods. Always*
wear a pair of latex gloves over your hands and fit nitrile gloves over the black rubber gloves inside the hood (Figure A.1).

Each dip should be recorded the dip in the Dipping Log. The time recorded for the dip should be the time the device is placed on the rotisserie. There should be two hours between each dip.

Figure A.1: Dipping mold, shown as separate pieces.

Proximal end – the end of the device with the channels

Distal end – the end of the device with the cap screwed on (the end w/o channels)
Day 1 – Dipping the Proximal End of the Device (All Pieces Separate)

1. Heat dipping hood (Figure A.3) up to 120 degrees Fahrenheit with a heat gun or another heating element. (It is easiest to set the timer the night before to start the heat gun a minimum of two hours at the 7.5 setting before the first dip. The green timer clip turns on the timer and the red clip turns the timer off.)
2. Clean mold (A.1) with rubbing alcohol.

3. Attach the end cap to the center piece of the mold.

4. Screw in gray blocks to the manifolds (side pieces).

Figure A.3: Dipping hood
5. Attach the aluminum bar to the distal end of the center piece of the mold
6. Push the long rotisserie wire through the hole in the steel bar and secure it with the pin (cylinder with small holes in it). Attach the other end of the wire to the hook on the rotisserie.
7. Start the rotisserie in order to lift the center mold (keep rotisserie wire taught so that it doesn’t get caught in the rotisserie).
8. Remove the lid from the BioSpan vat and place the vat under the device.
9. Reverse the rotisserie direction and slowly drop the mold into the BioSpan.
10. Stop the mold once the BioSpan has reached past the small steel pins (or just past the proximal critical area).
11. Reverse the rotisserie to remove the mold from the BioSpan. When the mold comes out of the BioSpan, immediately start a timer for four minutes as you let excess BioSpan drip off. (It is unnecessary to leave the mold in the BioSpan for a

Figure A.4: Gray blocks attached to each housing manifold
certain period of time. Once it is dipped, it can be removed immediately afterwards.)

12. Once four minutes are up, insert the longest screw with a nut glued on to the proximal end of the centerpiece.

13. Lower the centerpiece and disconnect the rotisserie wire from the aluminum bar and from the rotisserie.

14. Secure the steel bar and mold to the rotisserie with the pin and clip. Loop the hanging wire around the screw at the proximal end (Figure A.5).

15. Dip the proximal side of one manifold (channel end) barely past the critical area.

16. Hold the manifold (channel side down) over the BioSpan and allow it to drip for four minutes.

Figure A.5: Rotisserie setup with the centerpiece and two side manifolds attached to the aluminum bar.
17. Attach the manifolds to the steel bar using the screws attached to the gray blocks and secure them with butterfly nuts.

18. Check for bubbles after five minutes. If there are bubbles, use the dentist tools or something sharp to pop the bubbles. Do not stop the rotisserie when popping bubbles.

19. Wait two hours.

20. Remove steel bar from rotisserie and repeat steps 4 – 15 one time. Remember to remove the long screw before dipping the centerpiece again.

21. Leave device on rotisserie overnight.

Day 2 – Dipping the Proximal End of the Device (All Pieces Assembled)

1. Stop rotisserie and remove device attached to aluminum bar from the hood.

2. Unscrew both manifolds from the aluminum bar and remove grey blocks.

3. Unscrew centerpiece from aluminum bar.

4. Using a scalpel, remove thin layer of BioSpan on both manifolds and centerpiece that cured in the attachment areas.

5. Unscrew and remove distal cap from centerpiece.

6. Wipe down all pieces with rubbing alcohol.

7. Slide both manifolds into the appropriate positions on the centerpiece, followed by the distal cap. Screw together distal cap and centerpiece to complete assembly (Figure A.2). (Try to minimize any gaps between the manifolds and the centerpiece. These make demolding difficult.)

8. Reattach the aluminum bar to the device through the holes in the distal cap.
9. Place device back in dipping hood and wait for the hood to reheat to 120 degrees Fahrenheit.

10. Push the long rotisserie wire through the hole in the steel bar and secure it with the pin (cylinder with small holes in it). Attach the other end of the wire to the hook on the rotisserie.

11. Start the rotisserie in order to lift the center mold (keep rotisserie wire taught so that it doesn’t get caught in the rotisserie).

12. Remove the long screw from the proximal end of the centerpiece (if it is attached).

![Figure A.6: Complete device, hanging from aluminum bar, ready to be dipped in Biospan.](image)

13. Remove the lid from the BioSpan vat and place the vat under the device.

14. Reverse the rotisserie direction and slowly drop the mold into the BioSpan.
15. Lower the device into the BioSpan until it is submerged to a level just past the proximal critical area.

16. Slowly raise the device from the BioSpan. Once completely removed, stop the rotisserie and allow it to drip while hanging for 5 minutes.

17. Place lid on BioSpan vat and set it aside.

18. Screw the long screw into the proximal side of the centerpiece.

19. Lower the device and disconnect the rotisserie wire from the aluminum bar and from the rotisserie.

20. Secure the steel bar and mold to the rotisserie with the pin and clip. Loop the hanging wire around the screw at the proximal end (to ensure a level rotation on the rotisserie).

21. Start the rotisserie at the slowest speed.

22. Check for bubbles after five minutes. If there are bubbles, use the dentist tools or something sharp to pop the bubbles. Do not stop the rotisserie when popping bubbles.

23. Let the device cure on the rotisserie for two hours and then repeat steps 10-21 two additional times. After the third dip, let the device cure on the rotisserie overnight.

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*Day 3 – Dipping the Distal End of the Device*

1. Stop the rotisserie and remove the long screw from the centerpiece.

2. Insert a short screw with nuts to both manifolds on their proximal ends.
3. Fit screws through holes on the cross-hook and secure the cross hook with wing nuts on each screw.

4. Remove the device and aluminum bar from the rotisserie.

5. Hook rotisserie wire onto the cross-hook. The other end of the wire should be attached to the hook on the rotisserie.

6. Raise the device up to a level above the vat height.

7. Remove the aluminum bar while the device is hanging.

8. Remove the lid from the BioSpan vat and place the vat under the hanging device.

9. Slowly lower the device using the rotisserie into the BioSpan vat.

10. Submerge the device in BioSpan to a level just below the proximal critical area.

11. Raise the device from the BioSpan and allow excess BioSpan to drip off of the device for 5 minutes.

12. Place lid on BioSpan vat and set the vat aside.

13. Re-screw the aluminum bar into the distal cap of the device while it is still hanging. (Make sure to keep pressure on the bar while screwing it in to make sure it is tightly secured. It is okay to feel around for the holes with the screws and touch the BioSpan at the bottom of the mold since you will be unable to see the holes at this point.)

14. Lower the device and disconnect the rotisserie wire from the aluminum bar and from the rotisserie.

15. Secure the steel bar and mold to the rotisserie with the pin and clip. Loop the hanging wire around the cross hook at the proximal end (to ensure a level rotation on the rotisserie).
16. Start the rotisserie at the slowest speed.

17. Check for bubbles after five minutes. If there are bubbles, use the dentist tools or something sharp to pop the bubbles. Do not stop the rotisserie when popping bubbles.

18. Allow the BioSpan to cure while the device is on the rotisserie for 2 hours.

19. Following 2 hours, repeat steps 4-25 four more times. On the final (fifth) dip, allow the BioSpan to cure while the device is on the rotisserie overnight.

Day 4 – Laying the Bead

1. Shut off the heat gun.

2. Remove the device from the dipping hood.

3. Unscrew all attachments from the device (aluminum bar, short screws, and cross-hook).

4. Wipe down the entire device with rubbing alcohol.

5. Collect a sample of BioSpan in a 3 mL syringe, attach an 18-gauge needle, and place in the fume hood.

6. Using the syringe, lay a thick line of BioSpan (lay the bead) along one proximal critical area, at the attachment area between the manifold and centerpiece (Figure A.7).
7. Place the device in the 120 degree Fahrenheit oven located in the fume hood for 2 hours to allow the BioSpan to cure.

8. Repeat step 6 on the other proximal critical area and place the device in the 120 degree Fahrenheit oven located in the fume hood overnight to allow the BioSpan to cure.

**Day 5 – Water Submersion**

1. Remove the device from the oven.

2. Completely submerge the device in a bucket of water overnight to remove any excess DMAC within the BioSpan.

**Day 6 – De-molding**

1. Remove the device from water and dry it off.
2. Cut away the BioSpan from the proximal and distal end faces of the device (both on the center piece and on the manifolds).

3. Carefully peel away the housing from the mold. Use alcohol as a lubricant to help prevent tearing and ripping. Slide a plastic paint mixer between the mold and the housing to help separate both components.

4. Once the housing is completely separated from the surface of the mold, remove the distal end cap from the device.

5. Slide the centerpiece out of the housing through the proximal end of the housing.

6. Remove both manifolds from the housing by pulling them through the distal end of the housing. The housing is now completely separate from the mold and the dipping process is complete (A.8).

![Figure A.8: cTAL housing, removed from dipping mold.](image)

**Leak Testing and Patching: Housing**

*Leak Testing Housing*

1. Place a distal cap (part of the mold), that is siliconed closed, in the bottom of the housing.
2. Use medical tape to tape the housing to the cap.

3. Fold over the top of the housing and use plastic clamps to clamp shut.

4. Attach the connectors to the manifolds and zip tie in place.

5. Fill device with water from the sink. Ensure there are no air bubbles in the device.

6. Clamp shut the tubing attached to the connectors when the device and connectors are completely full of water.

7. When the device is completely full of water attach a pressure transducer to one of the lure locks and ensure the other lure is closed.

8. Calibrate the transducer using the hand pump (at 0 and 300 mmHg).

9. Fill the pressure transducer and line with water

10. Take the pressure using the Biopac system

11. Check for leaks, if the device is leaking mark the leak then drain the water out of the device by releasing the clamps

12. If the pressure is below 60 mmHg add more water using a 60 cc lure locking syringe

13. Repeat steps 14-16 until the pressure is 60 mmHg

14. If the device has no leaks at 60 mmHg then drain the water by releasing the clamps

15. Remove all of the tape and attachments

16. Wipe dry with paper towel

17. Let housing dry for 15 min before patching.
Patching Housing with Biospan

1. Wipe down the area to be patched with alcohol and allow it to dry.
2. Use a syringe to pull 1-2 ml of BioSpan from a jar of 24% BioSpan and place it and the device in the oven (at 120° F) for 10 minutes to allow it to heat up.
3. If necessary, place a piece of tape on the underside of the housing to prevent BioSpan from seeping through while patching/curing.
4. Attach a needle to the syringe and use this to apply a bead of BioSpan to the hole in the housing.
5. Hold the housing in place for almost 15 minutes until the BioSpan sets and won’t move once you place it into the oven.
6. Be careful with the BioSpan! If not used properly, the BioSpan will start to eat away at the existing BioSpan – so once you place the bead onto the housing, don’t wipe it away again!
7. Leave device in the oven overnight and the device will be ready to leak test the next day.

Rolling Fiber Bundles

Making the fiber bundle for the compliant device is a relatively straightforward process. One of the most important precautions to take when wrapping the fiber is to make sure everything is sterile and stays sterile. Any unnecessary particles in the fiber have the potential to impede flow through the device. Before beginning, make sure the bench top
is clean and the blue pads are replaced if necessary. A lint roller is helpful to clean these off if any loose particles are on top. Wear gloves when wrapping and touching the fiber.

Making the Core for the Fiber Bundle

1. See Purpose.

2. From the dimensioned drawing in Figure A.9, sketch the outline of the core on a piece of polycarbonate (thickness = 1/16”). Note that the drawing found in this document is not to scale, so the design must be redrawn on the polycarbonate, not simply traced.

![Figure A.9: Polycarbonate core for the cTAL fiber bundle.](image)

3. Using the dremel (with the heavy-duty cutoff wheel attachment) located under the bench in a toolbox, cut out the core from the sheet of polycarbonate. It is
important to wear safety glasses during this procedure as bits of material may fly off.

4. Remove the backing on the polycarbonate

5. Use a piece of sandpaper to sand off any rigid marks along the sides of the core. The edges of the core should be smooth so they do not tear the fiber when wrapping.

6. After sanding the edges, wash the core off in the sink with water and a little bit of rubbing alcohol.

7. Clean up the polycarbonate fragments and work area.

Wrapping the Fiber

1. See Purpose.

2. Obtain a completed core (H bracket) and wash it off with water and rubbing alcohol.

3. Locate a ring stand with a long rod attached to it and then place the white roll of fiber on the end of the rod.

4. Pull out a length of fiber about 16 inches long.

5. Locate the double-sided tape in one of the drawers and put a piece of this tape on one side of the long end of the H bracket. The other side of the tape will be placed on the end of the fiber strand you pulled out.

6. Flip the H bracket over and attach the end with tape on it to the fiber end as close as possible to the melted edge of fiber. Make sure to place the core vertically centered on the fiber.
7. Trim off any extra fiber if needed before you start wrapping.

8. Now you should have the H Bracket attached and laying on top of the fiber. You have to wrap the fiber around the core **42 times for the cTAL**. You can count each complete revolution around the initial point of attachment as 1 wrap, or you can count out **84 flips**. You already have 1 flip if you have the H bracket attached to the fiber correctly.

9. You must make sure that the fiber is even on both ends when you are wrapping it. There are two layers of fiber and they **must be kept even** when wrapping. One layer should not be higher than the other for example.

10. When wrapping, do not wrap with tension on the fiber. You should pull out a strand and flip the bracket over freely. Do not wind the fiber tightly, but also make sure the fiber is not loosely wound to the core.

11. Remember to stay sterile throughout the wrapping procedure.

12. Fiber bundles should be 3.6 ± 0.1 cm thick when finished.

![Rolled fiber bundle with thickness of 3.6 cm.](image)

Figure A.10: Rolled fiber bundle with thickness of 3.6 cm.
13. Once you have completed the wrapping, locate the Impulse Sealer on the shelf of the bench top. It is long and blue. Plug it in and turn the heating knob to 5.

14. This is used to melt the two layers of fiber into one. It is very easy to use and all you have to do is plug it in and turn the heat knob on the right to about 4 or 5. Let it heat for about thirty seconds.

15. Now place the completed fiber bundle on the left of the Sealer so you have enough fiber on the right to fold up on the side of the fiber bundle without impeding flow through the top of the bundle. If you melt the fiber too far to the right, when you fold it up, the tape will be attached on the top side of the bundle, ultimately impeding flow through the bundle.

16. Push the black handle of the Sealer down to melt the fiber for about 5 seconds and release.

17. Pull about 1 inch of fiber through the sealer to melt the fiber in a second place.

18. You should now have two distinct melt “lines” on the fiber.

19. Cut closely down the edge of the first melt line closest to the wrapped bundle.

20. Now you need to attach the end of the fiber to the side of the bundle. Cut a piece of double sided tape, put it on the side of the fiber bundle, and fold the end of the fiber up, attaching it securely.

21. Now you have to melt the ends of the fiber bundle using a Hot Plate. It is located on the first shelf. Plug it in and turn the heating knob to 5. It takes a couple minutes to warm up. There should be a small, scrap plate of aluminum on top of the Hot Plate. Leave this plate on the Hot Plate and make contact with this as
opposed to the Hot Plate itself when melting. This is solely for the purpose of not accruing melted debris on the Hot Plate.

22. You need to melt the outer 10 layers or so of fiber together. Do not place the end of the bundle on the hot plate too long or you will melt too much of the fiber.

Melt the other side of the fiber bundle.

23. Once you have melted both ends of the fiber, check for any loose particles on the bundle and remove them. Locate a completed housing and carefully insert the fiber bundle through the bottom of the cured compliant housing.

24. Put the completed housing with fiber bundle inserted into a large plastic bag and into the box labeled “Complete Housings and Bundles”.

Potting: Binding the cTAL fiber bundle and housing

Preparing Devices for Potting

1. Fiber bundles should be 3.6 ± 0.1 cm thick.

2. Insert bundles into housing – try not to pull on fibers, try to keep them aligned.

3. Make sure the bundles extend a minimum of .75 inches from the critical area on each side of the housing.

4. Trim the housing down on the proximal side until the housing no longer extends past the bundle.

5. Fold back channels and using painter’s tape, tape them down.
6. Measure out one 6” x 24” piece of saran wrap and wrap around device. Use painter’s tape to tape down the border of the saran wrap, keeping 0.75” of the device exposed on each end.

7. Cut a triangle on both the proximal and distal ends of the device, and on the same edge. It should be large enough that you can’t see any of the housing when looking through the holes in the top of the potting caps.

**Preparing Centrifuge**

1. Wipe down potting caps with alcohol and use paper towel to grease with Grease-It.

2. Spray all threaded rods with Grease-It under the hood.

3. Assemble cages around devices. Before tightening screws, make sure the device is touching neither the top nor the bottom of the potting caps. Insert a micro-spatula into top and bottom of potting caps to make sure there is sufficient room. Also, make sure housing is in potting caps and is not folded back.

4. Hand tighten screws first, and make sure the potting caps haven’t warped when tightening with a wrench.

5. Take four elbow connectors and two straight connectors and trim them using a dremel so that they do not extend past the inner rims of the potting caps and the potting tray respectively.

6. Duct tape the rim on the table where the centrifuge hood meets the table (Figure A.11).
Preparing Potting Material

At all times during preparation, keep materials A and B separate!

1. Prepare cartridges by marking them with appropriate 5, 135, and 145 mL lines (see the master for measurements).

2. Duct tape the bottom (smaller) opening close.

3. Use 20 oz syringes to pour A and B into their respective cartridges up to the 145 mL line.

4. Label each cartridges as A and B.

5. Cap the remaining open side with the given black cap.
6. Tip the cartridge over when the cap is half way closed and vent the duct taped side so that pressure does not build up inside the cartridge.

7. Snap cartridges A and B together.

8. Load the cartridges into the gun.

9. Open the tape again to allow for venting and pump the lever until the material fills the cartridge to the 145 mL line.
   a. Make sure there is an equal amount of potting material in each cartridge.
   b. If material spills out of the cartridges, allow the tape to act as a barrier and DON’T LET A AND B MIX OR THEY WILL START TO SOLIDIFY.

10. Press the silver lever and pull back on the handle to release the pressure on the cartridges.

11. Take the cartridges out and replace the duct tape on each.

12. Keep the cartridges in the hood when not using them.

Potting Procedure

1. Pull down the lid of the centrifuge and make sure it is locked. Go through the safety checklist.
2. Turn on heating element on centrifuge and turn display on centrifuge to 13
(corresponding to 224 RPM). Whenever increasing or decreasing speed on
centrifuge, adjust by increments of three, wait for centrifuge to reach speed and
then repeat.

**SOMEONE SHOULD BE WATCHING THE CENTRIFUGE AT ALL TIMES. BE READY TO HIT THE EMERGENCY STOP IF SOMETHING GOES WRONG.**

3. Place both cartridges in oven for half an hour with black plungers facing down
and tape facing up.

4. Speed centrifuge up to 41 on the display (707 RPM) after 30 minutes.

5. While speeding up centrifuge, attach the spiral mixing nozzle to the cartridges and
then the black threaded cap on top. The black cap will not fit all the way, just
rotate as far as possible.
6. Put cartridges in caulk gun and purge 10 ml of potting material into waste tray.

7. Once 707 RPM is reached, open cover from hole in the top of the centrifuge and inject 130 ml of potting material evenly into the middle of the white tray. Stop with 5 ml of potting material in each cartridge. Do not allow potting material to accumulate past the holes in the side of the tray.

8. Close cover and allow centrifuge to spin for twelve minutes at 707 RPM.

9. Purge the last 5 ml into the waste tray.

10. After twelve minutes are over, decrease speed to 13 on display. Let centrifuge run for 48 minutes.

11. Decrease speed to 5 on display. Once it reaches 5, hit stop button. Once centrifuge has completely stopped rotating, hit both emergency stop buttons.

12. Turn off heat gun, unplug centrifuge, and allow cooling down for 10-15 minutes.

13. Plug centrifuge back in and lift hood a few inches. Let centrifuge vent for an hour.

14. Separate cages from centrifuge and reattach facing the opposite direction.

15. Prepare potting material and repeat potting procedure on unpotted side.

**Cutting: Exposing the ends of the fibers**

After removing the devices from the potting setup, the potted ends must be cut to open up and expose the hollow fibers.

1. Measure ¼” in from the inside edge of the potting (on both sides) and mark with a sharpie. This is how far in you will cut (1/4” of potting must remain on either side of the cTAL after you are finished). Figure A.13.
2. Tape back the inlet and outlet manifolds so they are not in the way for cutting.

3. Place the cTAL in the cutting fixture so that the potted ends are exposed, up to the sharpie marks. Tighten down all screws (Figure A.14).

4. Make sharpie marks at the exposed potting – cutting fixture interface. This will be your gage as to whether the cTAL is moving in the fixture during cutting.
5. Use the cutting apparatus at MC3 (Figure A.15). Bring along a sharpie, ruler, hex key (for screws), and alcohol.

![Figure A.15: Cutting apparatus](image)

6. Wear the protective gloves when using the cutting apparatus.

7. Adjust the blade stop/arm on the cutting bench so that the blade can pass through the cTAL without hitting the cutting fixture.

8. Align the cTAL/cutting fixture with the blade. Squirt alcohol on the blade.

9. Begin cutting. Take the smallest slices possible. Reapply alcohol to the blade as necessary.

10. IMPORTANT: Make sure the cTAL is not being pulled from the cutting fixture. Watch your sharpie marks closely so you don’t cut too far into the potting.

11. Cut up until the sharpie marks.

12. Switch sides and repeat.

13. Remove cTAL from the cutting fixture and repeat with another device.
Leak Testing and Patching: Potted and Cut Devices

Leak Testing Cut Devices:

1. Attach connectors to manifolds and zip tie in place
2. Fill device with water from the sink.
3. When the device is completely full of water attach a pressure transducer to one of the lure locks and ensure the other lure is closed.
4. Fill the pressure transducer and line with water and calibrate the transducer using the hand pump.
5. Take the pressure using the Biopac system
6. Check for leaks in both the housing and potting material, if the device is leaking mark the leak with a permanent marker, then drain the water out of the device by releasing the clamps
7. If the pressure is below 60 mmHg add more water using a 60 cc lure locking syringe
8. Repeat steps 8 and 10 until the pressure is 60 mmHg
9. If the device has no leaks at 60 mmHg then drain the water by releasing the clamps
10. Remove all of the connectors
11. Dry out device and fiber bundle using filtered air for at least 4 hours before patching.
Patching Cut Devices: Housing

1. Ensure the housing or potted device is dry before patching.

2. Using a 3 cc lure lock syringe, draw up 1 mL of BioSpan from the vat in the Dipping hood and attach an 18 gauge needle to the syringe. MAKE SURE THE DIPPING HOOD IS CLOSED WHILE THE BIOSPAN VAT IS OPEN.

3. If necessary, tape down manifolds to keep them out of the way of the patched holes.

4. Place device and syringe in 120 degree farenheight oven for 10 minutes to heat up.

5. Only patch one side of the device at a time to prevent BioSpan from dripping or spreading to unwanted areas.

6. To patch holes, dap a small amount of BioSpan onto the hole.

7. To patch slits, tape along the slit on the inside of the device. Then lay a thin line of BioSpan along the slit.

8. Place device and syringe back in oven and let it cure for at lease 3 hours before patching the other side and at least 12 hours before leak testing again. The patched side of the device should be facing up.

Patching Cut Devices: Potting

1. Ensure the device is dry before patching.

2. Clean out 3, 1 mL syringes with alcohol to remove lubricant.

3. Draw up 1 mL of potting material Part A and 1 mL of potting material Part B.

4. Eject both parts of the potting material into a weigh dish and mix together.
5. Using either the third syringe or a popsicle stick, lightly dab/eject a little bit of potting material onto any holes in the potting.

6. Place weigh dish and device into the 120 degree farenheight oven (patched side facing up). Let the potting material dry for at least 2 hours before patching the other side, or overnight before leak testing again.

**Attaching Gas Caps and Performing Air Leak Tests**

1. Epoxy connectors in the acrylic gas caps

2. Fit the gas caps to the cut ends of the cTAL. Use sandpaper and/or the sanding attachment on the Dremel to make the caps fit properly.

3. Apply silicone around the edge of the cut end and then insert into the gas cap.

4. Press firmly in place

5. Apply silicone at the gas cap – cTAL edge/interface so that it is completely sealed.

6. After attaching both caps, set aside and allow it to cure for 24 hours.

Figure A.16: cTAL with gas caps
7. Fill a reservoir, large enough to fit the cTAL, with water.

8. Clamp the inlet and outlet of the cTAL with blue clamps so no air can escape.

9. Attach tubing to the 1.4” connectors on the gas caps

10. Attach the tubing to air and blow air (lightly) through the device while the device is submerged under water.

11. Watch for air leaks from the gas caps.

12. If there are any leaks, mark them with a sharpie and then patch them with silicone after the outside of the device is dry.

13. Allow to cure again for 24 hours and repeat gas leak testing.

14. Repeat until no air leaks remain.