Proactive esophageal cooling protects against thermal insults during high-power short-duration radiofrequency cardiac ablation

Marcela Mercado Montoya, Tatiana Gomez Bustamante, Enrique Berjano, Steven R. Mickelsen, James D. Daniels, Pablo Hernandez Arango, Jay Schieber, and Erik Kulstad

Proactive esophageal cooling with a novel cooling device has been shown to reduce endoscopically identified thermal injury during radiofrequency (RF) ablation for the treatment of atrial fibrillation using medium power settings. We aimed to evaluate the effects of proactive cooling during high-power short-duration (HPSD) ablation. A computer model accounting for the left atrium (1.5 mm thickness) and esophagus including the active cooling device was created. We used the Arrhenius equation to estimate the esophageal thermal damage during 50 W/10 s and 90 W/4 s RF ablations. With proactive esophageal cooling in place, temperatures in the esophageal tissue were significantly reduced from control conditions without cooling, and the resulting percentage of damage to the esophageal wall was reduced around 50%, restricting damage to the esophagus and consequently sparing the remainder of the esophageal tissue, including the mucosal surface. Lesions in the atrial wall remained transmural despite cooling, and maximum width barely changed (<0.8 mm).

Conclusions: Proactive esophageal cooling significantly reduces temperatures and the resulting fraction of damage in the esophagus during HPSD ablation. These findings offer a mechanistic rationale explaining the high degree of safety encountered to date using proactive esophageal cooling, and further underscore the fact that temperature monitoring is inadequate to avoid thermal damage to the esophagus.

1. Introduction

Atrioesophageal fistula (AEF) is a serious complication of ablation procedures during the performance of pulmonary vein isolation (PVI) for the treatment of atrial fibrillation, with a mortality rate as high as 80% [1,2]. Although the rate of AEF after PVI is often assumed to be around 0.2%, AEF is most certainly underreported. This is due to the fact that the presentation of AEF is delayed, the clinical picture mimics the much more common presentation to the emergency department of sepsis, and the diagnosis is difficult even if the condition is suspected [3,4]. Recent randomized controlled trials also suggest the rate may be higher than generally appreciated [5–7]. Various approaches have been taken to reduce the occurrence of AEF, including reducing power to the posterior wall, monitoring luminal esophageal temperature (LET), deviating the esophagus, and actively cooling the esophagus. Many of these approaches have been shown to be either ineffective or to potentially increase esophageal injury rates [8–13].

Active cooling can be performed either reactively, with manual cold-water instilled via gastric tube in response to measured LET rise, or proactively, with a dedicated cooling device providing cooling throughout the duration of the ablation procedure. Reactive cooling (where cooling is provided after a thermal insult has been detected by a LET monitoring probe) has been shown to have some benefit in a meta-analysis in reducing severe esophageal injury by 61% [14]. On the other hand, proactive cooling (where the temperature of the esophagus is preemptively reduced before ablation begins, using a dedicated cooling device) has shown greater benefits, with reductions in severe esophageal lesion formation of 67%–83% [15–17]. The rapidly growing use of proactive esophageal cooling during PVI has resulted in the publication or presentation of data on thousands of patients, and well over 10,000 ablations have now been completed with no AEF formation yet reported, and only a single peri-cardio-esophageal fistula known to have occurred [18–20].

With better-than-expected clinical safety being reported using proactive esophageal cooling, additional mechanisms of protection beyond mere dissipation of heat appear to be involved. A thermal insult is almost certainly a prerequisite for AEF formation [21,22], with thermal latency contributing to lesion growth even after the cessation of radiofrequency...
(RF) energy application [23,24]. Recent developments in the field of burn injury healing suggest that alteration of cellular behavior with cooling may also be a factor in preventing injury progression (by decreasing release of lactate and histamine, stabilizing thromboxane and prostaglandin levels, altering membrane permeability, inhibiting kallikrein activity and altering gene expression) [21,25–28]. With attainment of lethal isotherm temperature in esophageal tissue likely the inciting event from which AEF development begins, we aimed to evaluate using computer modeling the effects of proactive esophageal cooling on thermal damage in the esophagus during high-power short-duration (HPSD) ablations.

2. Methods
We used mathematical modeling to study the physical phenomena involved during RF catheter ablation of the left atrium with and without proactive esophageal cooling. A 3D model was built and solved using the finite element method with the software COMSOL Multiphysics (COMSOL, Burlington, MA, USA). The model makes use of Pennes’ Bioheat equation and includes accommodations for the phase change of tissues during the RF ablation.

2.1. Model geometry
Figure 1(a) shows the active cooling device on the left panel, and Figure 1(b) shows the model geometry, including all the relevant tissues and their proximity to the ablation target: atrial wall, epicardial fat, esophagus/connective tissue and blood (cardiac chamber). A cylindrical structure was assumed to be embedded into the connective tissue in order to model the esophageal lumen occupied by an active cooling device (ensoETM, Attune Medical, Chicago, IL, USA). This device was modeled as a hollow silicone tube (1.2 cm diameter, 0.65 mm wall thickness) full of cold water which is circulated by a pump. Control conditions, in which no active cooling device was present, were modeled by a collapsed esophagus. In this case, the subdomain corresponding to the cooling device was replaced with the properties of ‘esophagus and other tissues’. The model considered an irrigated-tip electrode (7Fr, 4 mm) placed perpendicularly on the atrial wall surface and inserted to a depth of 0.5 mm. Only half of the physical domain was considered as both the geometry and the physics are symmetric. Tissue thickness was set to average posterior-wall dimensions: 1.5 mm, 0.75 mm and 2 mm for the atrial wall, fat layer and the esophagus [29–31].

2.2. Governing equations
The Maxwell equations and the Bioheat equation were used to solve electric and thermal problems involved during RF ablation. The degree of thermal damage was calculated using the Arrhenius model [21,24,31,32]. In electromagnetic terms, the skin depth at 500kHz for biological tissues is on the order of 1 mm, suggesting that there are hardly any opposing eddy currents induced by the changing magnetic field resulting from the RF current. In other words, electric and magnetic fields can be considered decoupled and, consequently, the electric field \( \mathbf{E} \) (V/m) can be computed by using the frequency-domain electric Maxwell equations (Equations (1–3)) [33]

\[
\nabla \cdot \mathbf{J} = 0
\]

\[
\mathbf{J} = \sigma \mathbf{E} + j\omega \mathbf{D}
\]

\[
\mathbf{E} = -\nabla \Phi
\]

where \( \mathbf{J} \) is the current density (A/m²), \( \mathbf{D} \) is the electric displacement (C/m²), \( \Phi \) is the electric potential (V), \( \omega \) is the frequency (500 kHz for RF catheter ablation) and \( \sigma \) is the electric conductivity (S/m). The electrical problem is computed in all the subdomains shown in Figure 1(b): blood, myocardium, catheter tip and body, esophagus, surrounding tissues and cooling device (wall and circulating water).

The thermal problem was solved using the Bioheat equation [34]:

\[
\rho C_p \frac{\partial T}{\partial t} + \nabla \cdot (-k \nabla T) = Q_{\text{bio}} + Q_{\text{RF}}
\]

where \( T \) is the temperature (°C), \( \rho \) is the density (kg/m³), \( C_p \) is the specific heat capacity at constant pressure (J/kg K) and \( k \) is the thermal conductivity (W/m² K). The thermal problem is computed in all the subdomains except in the blood. Additionally, a term corresponding to thermal advection is considered specifically for the subdomain corresponding to the cold circulating water (i.e. inside the silicone tube), leaving Equation (4) as follows:

\[
\rho C_p \frac{\partial T}{\partial t} + \nabla \cdot (-k \nabla T) + \rho C_p \mathbf{u} \cdot \nabla T = Q_{\text{bio}} + Q_{\text{RF}}
\]

where \( \mathbf{u} \) is the velocity field for this subdomain (m/s), which is obtained from the inner area of the tube and the rate used in clinics (60L/h). The degree of thermal damage \( \alpha \) is described according to the Arrhenius formulation (Equation (6)) where the fraction of damage (FOD) \( \theta_d \) ranges from 0 and 1, and can be calculated with Equation (7).

\[
\frac{\partial \alpha}{\partial t} = A e^{-\frac{E_a}{RT}} (1 - \alpha)^n
\]

\[
\theta_d = \max \{0, \min(\alpha, 1)\}
\]

where \( A \) is the frequency factor (s⁻¹), \( E_a \) is the activation energy (J/mol), \( n \) is the reaction order and \( R \) is the gas constant (8.3145 J/mol-K). Values of 0.63 (63%) and 0.99 (99%) are commonly used to represent permanent tissue damage [24,35]. Lesion sizes were estimated at two times: just after the RF pulse, and up to 90 s after RF pulse onset to account for lesion expansion due to thermal latency. The heat sources \( Q_{\text{bio}} \) and \( Q_{\text{RF}} \) (W/m³) correspond to biological and electromagnetic phenomena, respectively. The term \( Q_{\text{bio}} \) is composed of metabolic (\( Q_m \)) and perfusion (\( Q_p \)):

\[
Q_{\text{bio}} = Q_m + Q_p
\]

The metabolic heat is negligible compared to the energy dissipation and was hence ignored [36]. The perfusion heat was given by:

\[
Q_p = \beta \rho_b c_{p,bl} \omega_{bl} (T_{bl} - T)
\]

where the following blood properties are involved: \( \rho_b \) is the density (kg/m³), \( c_{p,bl} \) is the heat capacity in (J/kg K), \( \omega_{bl} \) is
the perfusion rate (s\(^{-1}\)), \(T_{bl}\) is the blood temperature (37°C) and \(\beta\) is a non-dimensional parameter to simulate that blood perfusion ceases once tissue is completely destroyed by thermal necrosis (\(\beta = 1\) while fraction of thermal damage remains less than 99% and \(\beta = 0\) for 100% damage). The perfusion rate for each tissue is taken from the ITIS database [37]. The RF-induced heat source is due to Joule effect and is given by:

\[ Q_{RF} = \frac{1}{2} Re(J \cdot E) \]  

(10)

The irrigated electrode was modeled with a heat source with a mathematical structure similar to blood perfusion and given by Equation (9)

\[ Q_{electrode} = \rho_{H2O} C_{p,H2O} \omega_{H2O} (T_{H2O} - T) \]  

(11)

where the following saline water properties are involved: \(\rho_{H2O}\) is the density (kg/m\(^3\)), \(C_{p,H2O}\) heat capacity (J/kg-K), \(\omega_{H2O}\) catheter irrigation rate (s\(^{-1}\)) and \(T_{H2O}\) water temperature (assumed to be 25°C). This method mimics the cooling effect by saline circulating inside the electrode, and it has been broadly used in computer models of energy-based cooled applicators [35].

### 2.3. Boundary and initial conditions

We modeled constant power RF ablation by applying a power value at the interface between the catheter tip and body. Some electrical losses happen in the body beyond the zone around the ablation electrode, and this loss has been shown to be approximately 20% [38]. As such, only 80% of the value used in clinical practice was used for the simulation. Two HPSD settings were simulated: 50 W for 10 s and 90 W for 4 s. The active cooling device considered cold water at 4°C (inflow temperature) flowing at a rate of 60 L/h in the \(x\) direction (see Figure 1(b)). The water velocity \(u\) was directly defined in the heat transfer equation (see Equation (5)) and the transport of water flow was not therefore studied using momentum equations. The blood subdomain was exclusively considered in the electric problem. The thermal effect of the circulating blood on electrode and myocardium surface was considered with convective heat flux boundary conditions. The convective heat flux \(\dot{q}\) (in W/m\(^2\)) through the blood-electrode and blood–myocardium interfaces was given by:

\[ \dot{q} = h(T_{bl} - T) \]  

(12)

where \(h\) is the heat transfer coefficient (in W/m\(^2\)-K) and \(T_{bl}\) is the blood temperature (37°C). The values for \(h\) were 610 W/m\(^2\)-K and 3346 W/m\(^2\)-K for blood-electrode and blood–myocardium interfaces, respectively, which simulated ablation sites with low local blood flow (0.085 m/s), as in patients with chronic atrial fibrillation and dilated atria [39]. The symmetry boundary condition was used for the symmetry boundaries (see Figure 1(b)). The rest of the external boundaries were set with 0 V for the electric problem and as thermal insulation (no flux) for the thermal problem.

### 2.4. Material properties

Table 1 shows the thermal and electric properties for each material [31,32]. The parameters for the Arrhenius damage model were: \(n = 1\), \(A = 4.43 \times 10^{16} \text{ s}^{-1}\) for fat and \(7.39 \times 10^{38} \text{ s}^{-1}\) for the rest of the tissues, and \(E_a = 1.30 \times 10^5 \text{ J/mol}\) for fat and \(2.59 \times 10^5 \text{ J/mol}\) for the rest of the tissues [31,32,40]. The variation with temperature of the electric conductivity, density, thermal conductivity and the heat capacity were considered for all the tissues. These variations were represented using the following equations with the temperature in °C:

\[ \sigma = \sigma_{\text{ref}} \left[1.0 + 0.015 \cdot (T - 37.0)\right] \]  

(13)

\[ C_p = C_{p,\text{ref}} \left[1.0 + 0.0032 \cdot (T - 37.0)\right] \]  

(14)

\[ k = k_{\text{ref}} \left[1.0 + 1.20 \times 10^{-3} \cdot (T - 37.0)\right] \]  

(15)

\[ \rho = \frac{k_{\text{ref}}}{C_{p,\text{ref}}} \cdot \frac{1.474 \times 10^{-7} \cdot [1.0 + 3.39 \times 10^{-3} \cdot (T - 37.0)]}{1016 \text{ s}} \]  

(16)

The variations of density and thermal conductivity were taken from COMSOL predefined functions for human

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**Figure 1.** (a) Physical situation to be modeled with an active cooling device located in the esophageal lumen. (b) Model geometry including RF catheter, tissues near the ablation site and an active cooling device located in the esophageal lumen. The evaluation line (black line) for post-processing is shown across the ablated tissues, from tip of RF catheter to edge of active cooling device (with tissues including myocardium, fat and esophagus).
myocardium while the reference values were taken from the ITIS database [37]. The electric conductivity and the heat capacity were specified to increase linearly (Equations (13) and (14)) and then change near 100 °C to account for the water loss associated with the evaporation phenomenon [39]. Equation (14) describes the heat capacity of the tissue during heating below the boiling point of water, but above this value, the formation of bubbles can occur. The effect of this phase change on the temperature distribution in the tissue is therefore included. Between 30 °C and 110 °C, Equation (17) describes the vaporization latent heat of water (ΔH_vap^H_2O, kJ/kg). In this equation, a = -2.4911 kJ/kg°C and b = 2.5060 × 10^3 kJ/kg. 

\[ \Delta H_{vap}^{H_2O} = aT + b \] (17)

Since the exact description of the possible phase change of the water contained in the tissue due to its temperature increase during ablation is not part of the scope of this study, our approximation is to define a factor that governs the heat transfer from the sensible to the latent enthalpy, which is nothing more than the relationship between the vaporization latent heat of water, evaluated in a temperature range around the boiling point, and the specific heat of tissue. We considered the boiling point of water at 1.0 atm, i.e. 100 °C.

The factor is selected for the temperature range expected (without considering a detailed model of the phase change of the water contained in the tissue). The factor in this case is 555 in a temperature increment of ΔT = 1 °C or 138.75 for ΔT = 4 °C. Both factors showed similar behavior of the peak temperature against time, so the second factor was utilized to achieve easier convergence and reduce the computational cost. Electrical conductivity (σ) was also expected to be affected by evaporation since water loss induces a decrease in the tissue’s ability to conduct electrical current. Our model included this effect by drastically reducing the σ value (by four orders of magnitude) for temperatures above 100 °C [21].

### 2.5. Solver and verification

The computational domain was discretized using a hybrid mesh consisting of both tetrahedral and quadrilateral elements (Figure 2(a,b)). The mesh was refined around the electrode, where the changes in the dependent variables and material properties are expected to present the most representative changes. The ‘Mesh Control Faces’ option in COMSOL was used to refine the mesh around the electrode area to guarantee perfect symmetry of the mesh in the yz plane. The mesh element quality histograms with different criteria (skewness, volume vs. rate and growth rate) for the elements around the electrode are shown in Figure 2(c–e). A good mesh must have as best element quality as possible, and the more the histogram distribution tends to the right, the better is the mesh quality. The convergence test was done by using a finer mesh with a duplicated number of elements, keeping the proportion of elements in each region and the element quality. The results for fraction of damage, used to calculate the lesion depth and width, were used for convergence criterion. The results between the original and finer mesh remained within 2%.

In clinical terms, the device pre-cooling time (which is the time the proactive cooling device is performing inside the patient body before RF pulses begin) is around 5–10 min, which is enough to achieve the stationary state. In consequence, in order to consider this pre-cooling time, a stationary step was added (the results showed that 5 min was enough to reach a steady state in which the temperature varies less than 0.1 °C). Then, a second frequency-transient step takes the stationary results as initial values to run the ablation procedure, the frequency-transient option in COMSOL allows calculating the electric field in the frequency domain and the temperature as a time-dependent variable. The solution time was set as the ablation duration according to the different ablation setups (50 W or 90 W), while the time step was set as 0.02 times the ablation duration. Finally, a third step is added to calculate the temperature and tissue damage 1 min after the ablation catheter is turned off with a step of 1 s.

The segregated solver was set to solve the electric currents with the BiCGStab iterative method, the temperature with the PARDISO direct method and the Arrhenius equation with the MUMPS direct method. For the second and third steps, the high nonlinearities of the model given by the extreme changes in the material properties (electrical conductivity and heat capacity) due to the water evaporation around 100 °C, were controlled by adjusting the nonlinear method to the COMSOL option automatic highly nonlinear
The solver tolerance and time step were set as 'physics controlled' and established automatically by COMSOL. Simulations of both the control condition (no active cooling) and the experimental condition with active cooling were completed.

3. Results

Figure 3 shows the progress of the electrical variables during the RF pulse in the control case, i.e. without proactive esophageal cooling. As expected, the power remains almost constant throughout the RF pulse, despite impedance variations. The impedance shows an initial decrease and a slight increase just after the end of the pulse, which causes variations in the electrical current. The applied voltage is then suitably modulated in order to keep the power constant.

Figure 4 shows the temperature distributions in the tissue undergoing HPSD with (protection) and without (control) proactive esophageal cooling. The impact of esophageal cooling on the increase in temperature in the vicinity of the esophagus is clearly visible. The average temperature increase of the cooling water inside the active cooling device was 0.22°C in any case (computed at the outflow).

Figure 5 shows the temperature progress at different depths below the electrode. The temperature just at the tip of the RF electrode undergoes a slight increase until it reaches about 45–50°C. Possibly 1 mm away is the hottest point, which shows rapid increases in temperature up to 100°C. As we move away from the electrode, the temperature peak occurs later, even after the RF power has ceased. For the specific simulated distances, the temperature progress at 3 mm is highly affected by the presence or absence of esophageal cooling: while reaching temperatures of 50–55°C for the unprotected case, it remained around 40°C when esophageal cooling is used. When the protection is used, the temperature close to the esophageal lumen (4 mm depth) remains below 20°C.

Figure 6 shows the lesion shapes computed for 50 W/10s and 90 W/4s, with (protection) and without (control) proactive esophageal cooling. The lesion depths (computed with fraction of damage \( h_d = 63\% \) after 90 s) in the case without proactive esophageal cooling (control) were 3.5 mm for 50 W/10s and 3.1 mm for 90 W/4s, decreasing in the case of cooling: 2.9 mm for 50 W/10s and 2.7 mm for 90 W/4s. Note that the fraction of damage incurred by fat is lower than that of myocardial or esophageal tissue, which is a consequence of tissue parameters incorporated into the Arrhenius equation and reflects known resistance of adipocytes to thermal insult.

It is noticeable that without active cooling in place, the fraction of damage in the esophagus incorporates an important percentage of the esophageal thickness, whereas with proactive esophageal cooling, the damage is restricted to the epi-esophageal surface, sparing the remainder of the
esophageal tissue, including the mucosal surface. This can be also confirmed in Figure 7(a), where the percentage of esophageal lesion (using $l_d = 63\%$) is around 40–60% without protection (control case) and 25–30% with protection (computed 90 s after RF pulse). Regarding the impact of the active cooling of the esophagus on lesion formation in the atrial wall, as shown in Figure 6, 100% of myocardial lesions remained transmural, and as shown in Figure 7(b), cooling had very little impact on the maximum width ($< 0.8$ mm).

4. Discussion

4.1. Main findings

Despite the availability of clinical data demonstrating the efficacy and safety of proactive cooling to reduce endoscopically identified thermal injury [15,41,42] during RF catheter ablation, there is no specific information about the lesion size and temperature dynamics during and after the application of high-power short pulses in the case of esophageal proactive cooling. Since it is very complex to obtain this type of information from clinical studies, we built an in silico model and conducted computer simulations to analyze the impact of proactive cooling on temperature distributions, lesion size across the atrial wall and esophageal thermal damage after applying HPSD ablation. Although there are recent experimental data describing the temperature dynamics after these pulses and emphasizing the importance of thermal latency [43,44], none of them considered the presence of a proactive cooling probe. Our motivation was thus to acquire valuable information about these issues, which might provide a physical explanation for the current clinical data.

We report here the first description of temperature across the esophagus during HPSD RF ablation while proactive esophageal cooling is performed. Our model suggests that proactive esophageal cooling protects against thermal insults during HPSD ablation, and does this by precluding the attainment of tissue temperatures reaching or exceeding lethal hyperthermic doses at the esophageal mucosa and across the majority of the esophageal tissue. This in turn significantly reduces the fraction of damage sustained in the esophagus, and offers a mechanistic rationale for the lack of AEF development and the solitary pericardio-esophageal fistula reported, despite thousands of uses to date of proactive esophageal cooling during HPSD ablation.

Results from the model with no active cooling align well with the experimental data recently published. In terms of lesion size, we obtained depths of 3.45 mm for 50 W/10s and 3.05 mm for 90 W/4s and maximum widths of $< 0.7$ mm for 50 W/10s and $< 0.8$ mm for 90 W/4s. These values are in reasonable agreement with those reported by Nakagawa et al. [44] in the beating heart: 4.8 ± 0.9 mm deep and 8.2 ± 1.4 mm wide for 50 W/10s, and 3.6 ± 0.6 mm deep and 7.9 ± 1.2 mm wide for 90 W/4s. The limitations of the experimental models must be also taken into account, since there is an important discrepancy between the reported lesion sizes, possibly due to the different specific conditions of each experimental setup (contact force, temperature limit programmed in the RF generator, etc.). For instance, Leshem et al. [45] reported lesion depths ranging from 1.8 to 3.6 mm for 90 W/4s in the

Figure 3. Progress of electrical variables (power, impedance, current and voltage) during RF pulse using 50 W/10s (top) and 90 W/4s (bottom). Note that power is almost constant during the RF pulse, with variations less than 2 W.
right atrial wall depending on the location (postero-septal and postero-lateral), and widths around 6 mm. Regarding the progress of the temperature inside the tissue, our results reproduce well the thermal latency observed in experiments [43,44]. However, note that a perfect fit is not possible in terms of absolute values of temperature since there is a large gradient in the area near the RF electrode, where the temperature undergoes huge changes in just 1 mm of distance (see Figure 5).

Results from the model with proactive cooling showed lower tissue temperature values around the esophagus (see Figures 4 and 5). In particular, the peak temperature at 2 mm away from the esophagus lumen (which corresponds approximately with temperature at 3 mm deep) reached \( \sim 55 \, ^\circ\text{C} \) without protection and \( \sim 40 \, ^\circ\text{C} \) with protection. This difference (15 °C) is larger than what we observed in an experiment based on phantom agar using an esophageal balloon cooled at 5 °C (7.5 °C) [46], possibly due to the lower power applied in that case (constant temperature at 55 °C and power always below 50 W) compared to the powers applied here (50 and 90 W). The cooling effect around the esophagus translated into a lower percentage of damaged esophageal wall (see Figures 6 and 7(a)). These findings corroborate the data from clinical literature reporting significant safety benefits [18–20,47,48]. A survey of users of proactive esophageal cooling found that 35% utilized 40 W on the posterior wall, and 44% utilized 50 W power settings on the posterior wall [49]. Accordingly, a large percentage of cases performed with proactive esophageal cooling involve high power on the posterior wall. Our findings that proactive esophageal cooling reduces the peak temperatures sustained in esophageal tissue during HPSD ablation offer a theoretical basis to explain the clinical data to date. Our results showed an increment of 0.22 °C in the water temperature inside the device, which is slightly less than that found in clinical practice in elective surgical procedures, where a larger surface
area is exposed to heat transfer from the patient, and an increase of water temperature inside the device of up to 0.4 °C has been reported [50]. Nevertheless, both values are very small, which is physically reasonable due to the high volume of coolant flow in clinical use (over 60 L/h). Although cooling shows significant protective effects in the esophageal tissue adjacent to the cooling surface, the effect on atrial myocardium appears unlikely to be of clinical significance.

Figure 6. Lesion shapes (computed with fraction of damage $\theta_d = 63\%$ after 90 s with the Arrhenius model) for 50 W/10s and 90 W/4s, with (protection) and without (control) proactive esophageal cooling. Left sided images show the case after the RF pulse, right sided images show the case after 90 s.

Figure 7. Esophageal lesion transmurality (a) and maximum lesion width in the myocardial wall (b) computed with two fractions of damage $\theta_d$ (63% and 99%) just after the RF pulse and after 90 s, for 50 W/10s and 90 W/4s, and with (Protection) and without (Control) proactive esophageal cooling.
Transmurality of atrial lesions remained at 100% despite active cooling (see Figure 6), while differences in lesion width are possibly negligible (< 0.8 mm), especially when linear lesions are created by overlapping consecutive punctual lesions. It is also important to remember that a tradeoff between achieving overall smaller lesions (which are needed to minimize esophageal damage, e.g. using proactive cooling) and the need for additional ablation lesions to ensure complete/continuous ablation patterns. This issue could become relevant in case of very short distances between the proactively cooled esophagus and the RF electrode.

It is interesting to discuss the combined effect of electrode irrigation/surface cooling and esophageal protective cooling. The former aims to push the hot spots deeper into the tissue to achieve deeper and better developed lesions, while the latter aims to avoid high temperatures reached in the esophagus. Despite the fact that there are two opposite thermal mechanisms ‘pushing’ the hottest point in the opposite direction, nothing impedes the simultaneous use of irrigated electrodes and esophageal proactive cooling. In fact, in current use in the United States, Europe, Australia and Israel, essentially all operators use irrigated RF catheters with proactive esophageal cooling [15,20,42]. This phenomenon is well illustrated in the progress of temperature at 3 mm depth (see Figure 5). While in the control case (without esophageal cooling) the temperature increases due to the ‘hot spot push’ toward the esophagus, in the case of esophageal cooling, the ‘additional push of the hot spot’ toward the endocardium tends to compensate for the first push, managing to more or less stabilize the temperature during the time the RF pulse is applied (around \(40 ^\circ \text{C}\)).

One of the novel aspects of this study is the specific focus on HPSD ablation. In HPSD ablation, the reliability of LET monitoring is further compromised due to the velocity of temperature rise. A recent study further highlights the inability of LET monitoring to predict thermal injury in HPSD ablation [51]. Our findings in this study suggest that proactive esophageal cooling with a dedicated device is efficient enough to accommodate the rapid changes in temperature that occur with HPSD ablation. Although exact mechanisms behind formation of AEF remain uncertain, growing evidence in the burn literature strongly points to mechanisms triggered by an initial thermal insult, and a subsequent inflammatory cascade that progresses over days to weeks [25–27,52]. Moreover, abundant data demonstrate the ability of cooling to reduce or prevent the progression of burn injuries [27,52,53]. As such, proactive cooling may have a synergistic effect on reducing or preventing the formation of fistulas by (1) reducing peak temperatures that cause the initial thermal insult (thereby limiting the severity of the initiating injury), and (2) lessening the severity of the inflammatory cascade that stimulates progression of burn injury.

4.2. Limitations

Mathematical models provide valuable insight into physical phenomena but may not reflect all clinical findings due to the inherent variability in clinical practice and physiologic parameters. Nonetheless, the elucidation of general principles and ranges of expected findings are helpful in understanding mechanisms involved, and the availability of abundant clinical data and recent experimental data as comparison further strengthens the validity of this model. Debate continues over the exact mechanisms of AEF formation, and additional factors beyond thermal injury may be involved, as discussed above. Although the model described here quantifies the decrease in thermal insult possible with proactive esophageal cooling, it cannot account for downstream effects that may mitigate the production of inflammatory mediators that cause injury progression over days to weeks. Reducing this inflammatory effect may further increase protection against AEF. On the other hand, comparison to existing clinical data may be a suboptimal metric, considering the innumerable variables and dynamic factors involved in the pathophysiology of esophageal thermal injury and AEF. This may be of particular relevance in this condition where varied tissue properties between patients as well as differences in contact forces employed could impact the findings. At this point, considerations such as these, as well as the proximity of the esophagus to surrounding structures and its mobility, can only be accounted for via changes to geometric positioning in the model; however, this model serves as a robust system to further investigate the influence of these factors. Likewise, the impact of the periesophageal vasculature in dissipating heat, as well as potential microvascular effects on ischemic injury and other dynamic phenomena with substantial inter-patient and temporal variation are only partially considered in this model using the Bioheat equation.

In terms of mathematical modeling, the study has also some limitations. First, the blood flow and electrode irrigation were modeled in a simplified way without solving the fluid dynamics. Although this approach is less accurate for predicting surface width, it reasonably predicts lesion depth, and hence it is reasonable to study the thermal impact in deep points (e.g. in the esophagus). Second, mechanical deformation of the endocardial surface was not modeled, i.e. ‘sharp insertion’ of electrode was assumed. Although it is known that this approach overestimates the lesion size compared to the case with ‘elastic insertion’ [54], it is expected that this will affect equally with and without proactive thermal protection, and hence the overall conclusions will not change. And third, while the Arhenius model was used to estimate the thermal damage, we recognize that other methods could provide different results [55,56]. Likewise, since the mechanism of thermal insult is the same with and without proactive thermal protection, it is expected that the conclusions will be qualitatively the same in case of using a different method to estimate the thermal damage.

5. Conclusions

Proactive esophageal cooling significantly reduces temperatures and the resulting fraction of damage in the esophagus during HPSD ablation. Esophageal damage was restricted to the epi-esophageal region, sparing the remainder of the
esophageal tissue, including the mucosal surface. These findings may offer a mechanistic rationale explaining the high degree of safety encountered to date using proactive esophageal cooling. Moreover, these findings further underscore the fact that temperature monitoring is inadequate to avoid thermal damage to the esophagus.

Disclosure statement

M.M.M.: employment with Insilico SE, consulting for Attune Medical; S.R.M.: employment with Acutus Medical, consulting for Attune Medical; T.G.B.: employment with Insilico SE; P.H.A.: employment with Insilico SE; E.K.: equity and employment in Attune Medical. The rest of the authors declare no conflict of interest.

Funding

Research reported in this publication was supported by the National Heart, Lung, And Blood Institute of the National Institutes of Health under Award Number R44HL158375 (the content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health) and by the Spanish Ministerio de Ciencia, Innovación y Universidades/Agencia Estatal de Investigación (MCIN/AEI/10.13039/501100011033 under grant RTI2018-094357-B-C21).

ORCID

Marcela Mercado Montoya http://orcid.org/0000-0002-1453-4690
Tatiana Gomez Bustamante http://orcid.org/0000-0002-3947-1390
Enrique Berjano http://orcid.org/0000-0002-3247-2665
Steven R. Mickelsen http://orcid.org/0000-0003-4219-8230
James D. Daniels http://orcid.org/0000-0003-3360-6175
Pablo Hernandez Arango http://orcid.org/0000-0001-6383-7770
Jay Schiebert http://orcid.org/0000-0002-9644-5236
Erik Kulstad http://orcid.org/0000-0002-9331-8266

Data availability statement

The data underlying this article will be shared on reasonable request to the corresponding author.

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