Effect of Global Cardiac Ischemia on Human Ventricular Fibrillation: Insights from a Multi-scale Mechanistic Model of the Human Heart

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Abstract

Acute regional ischemia in the heart can lead to cardiac arrhythmias such as ventricular fibrillation (VF), which in turn compromise cardiac output and result in secondary global cardiac ischemia. The secondary ischemia may influence the underlying arrhythmia mechanism. A recent clinical study documents the effect of global cardiac ischemia on the mechanisms of VF. During 150 seconds of global ischemia the dominant frequency of activation decreased, while after reperfusion it increased rapidly. At the same time the complexity of epicardial excitation, measured as the number of epicardial phase singularity points, remained approximately constant during ischemia. Here we perform numerical studies based on these clinical data and propose explanations for the observed dynamics of the period and complexity of activation patterns. In particular, we study the effects on ischemia in pseudo-1D and 2D cardiac tissue models as well as in an anatomically accurate model of human heart ventricles. We demonstrate that the fall of dominant frequency in VF during secondary ischemia can be explained by an increase in extracellular potassium, while the increase during reperfusion is consistent with washout of potassium and continued activation of the ATP-dependent potassium channels. We also suggest that memory effects are responsible for the observed complexity dynamics. In addition, we present unpublished clinical results of individual patient recordings and propose a way of estimating extracellular potassium and activation of ATP-dependent potassium channels from these measurements.

Introduction

The heart is an electromechanical pump, where contraction is triggered and synchronized by electrical activation originating from the sinoatrial node. Abnormal initiation or conduction of the electrical impulses could result in a cardiac arrhythmia. Cardiac arrhythmias are an important cause of sudden and premature death in the industrialized world. In many cases the lethal event is ventricular fibrillation (VF). During VF, rapid and self-sustaining electrical activity in the ventricles acts to suppress the natural pacemaker, resulting in uncoordinated, weak and rapid contractions, which lead to death within several minutes [1].

VF often occurs as a result of acute regional cardiac ischemia, which is a condition when blood flow to part of the heart is substantially decreased, for example by reduced flow through a coronary artery [2]. In addition, global ischemia unavoidably accompanies VF, because the abrupt fall in cardiac output resulting from VF also results in compromised myocardial perfusion. Thus an episode of spontaneous VF will result in a progressively ischemic heart. The effect of this secondary ischemia on electrical activity during VF is important clinically, because defibrillation typically occurs several minutes after the onset of VF, and thus the mechanism is likely to have been modified by ischemia [3,4].

It is known that ischemia profoundly affects the electrophysiological properties of cardiac cells and tissue [5]. During VF, the rapidly changing patterns of electrical activity in the ventricles are sustained by re-entry, in which waves of electrical activation continually propagate into regions of recovered tissue [6]. Re-entry is seen as a spiral wave on the surface of the heart, and a scroll shaped activation wave in 3D cardiac tissue [7]. The question of how ischemia influences the behaviour of re-entrant activity during VF is important, and has been addressed by clinical, experimental and modeling studies.

Two important characteristics of VF are the frequency and spatiotemporal complexity of activation patterns, and these have been studied in animal heart experiments. Experiments on canine hearts [6] have shown that both activation rate and pattern...
Cardiac arrhythmias are an important cause of death in the industrialized world. The most dangerous type of cardiac arrhythmias is ventricular fibrillation. If left untreated, it leads to death within just few minutes. In most of the cases ventricular fibrillation occurs as a result of cardiac ischemia, which is a shortage of blood supply to the heart muscle. Furthermore ventricular fibrillation leads to decreased cardiac output, which in turn results in secondary ischemia. A recent clinical study investigated the effect of secondary ischemia on the organization of ventricular fibrillation in the human heart. However, in the clinical study it was not possible to obtain the whole picture of activation of the heart and to separate the relative roles of different processes induced by ischemia in the alterations to the cardiac electrical activity. In this study we use computer modeling to address these problems and to complement the results of the clinical study. Our results allow us to explain the change of electrical activation pattern in the heart during the first minutes of ischemia and to estimate the relative rates of those ischemia-induced physiological processes. We also present previously unpublished data on individual patient recordings from the clinical study.

In spite of the significance of these animal studies, the most valuable question for clinical practice is how global ischemia modulates the mechanism of VF in the human heart. Both clinical and modelling studies have established that the organization of VF in the human heart is quantitatively different to that in canine and porcine hearts [9, 10], with VF in the human heart shown to be characterized by a much lower VF complexity compared with animal hearts of similar size. For this reason, experimental studies on human hearts are extremely important for understanding underlying mechanisms. Recently this gap has partially been filled by studies in isolated myopathic hearts [11, 12], and studies in the in-situ human heart [13].

In the in-situ study [13], electrical activity was mapped on the heart surface during VF in ten patients undergoing routine cardiac surgery with cross-clamp fibrillation. Following the commencement of cardiopulmonary bypass to support the systemic circulation, the following protocol was used: (1) VF was induced by burst pacing, (2) after 30 s, global cardiac ischemia was initiated by applying an aortic cross-clamp, (3) after 2.5 min. ischemia, cardiac perfusion was restored by release of the cross-clamp, (4) recording continued for 30 s during reperfusion. During all 3.5 minutes of VF, electrical activity was recorded using 256 unipolar electrodes sewn into an elastics sock placed over the entire ventricular epicardium by the surgeon. Consistent with studies in animal hearts, the activation rate of fibrillation gradually decreased during global ischemia and increased abruptly after reperfusion. In contrast with animal hearts, the complexity of fibrillation patterns (measured as the number of phase singularities) continued to increase gradually during ischemia, although the data showed large variation.

The above mentioned study [13] had several important limitations, in common with other studies in human and animal hearts. First, electrical activity was recorded only on the epicardial surface, whereas electrical activation patterns in the ventricles are 3-dimensional. Second, ischemia is a complex process which involves several separate mechanisms including hyperkalemia, hypoxia and acidosis, and each of these components has its own time course in ischemia [5]. However, it is not generally possible to measure the relative contribution of these individual components to the excitation patterns in vivo.

Because it is difficult to measure all of the quantities of interest, even in animal hearts and tissue, detailed mechanistic and multi-scale models of cardiac electrophysiology at cell, tissue, and whole organ scales are becoming important research tools for the study of arrhythmia mechanisms. It was shown previously that the combination of modeling with experimental and clinical studies can provide valuable insights into arrhythmia mechanisms [9, 10, 14].

In this paper, we describe a detailed and comprehensive modeling study, which seeks to establish mechanisms that are consistent with the changes in activation patterns observed during VF in the human heart with global ischemia [13]. In addition, we present previously unpublished clinical results from [13], namely the changes in dominant frequency (DF) for individual patients. The model we use is a representation of human ventricular tissue, with cellular electrophysiology described by the TNXN06 model [15]. This model was adapted to enable us to investigate the effects on VF of three components of ischemia [5] both separately and in combination. Briefly the effects of these three components on the electrophysiological processes can be characterized as follows. Hyperkalemia is elevation of the extracellular potassium concentration, which leads to a shift of the resting potential of cardiac cells. Acidosis is an increase in intracellular pH, which results in smaller fast sodium and L-type calcium currents during depolarization. Hypoxia is a reduced oxygen supply, which impairs cellular metabolism and changes the ratio of ADP to ATP concentration inside the cell. In turn, this change results in the opening of specific ATP-dependent potassium channels. To describe them in our simulations, we developed a new model of the human K\textsubscript{ATP} channel based on experimental data from human cardiac cells [16].

We first determined the effect of each of the three components of ischemia on action potential duration (APD) and conduction velocity (CV) restitution in 1D models, along with the effects on re-entry in 2D models. Then we performed simulations of VF using a 3D anatomical model of the human ventricles [10], where the dependency of the activation rate of VF on each component of ischemia was determined.

Next, we used these dependencies to interpret the clinical recordings for individual patients. We showed that the predominant factor responsible for the change in activation rate during ischemia is hyperkalemia, and estimated the magnitude of this effect in each individual patient recording. Then we studied how the three ischemic components affect the complexity of VF expressed by the number of scroll wave filaments. Although there was a big variation, the results we obtained were qualitatively similar to the clinical recordings and we concluded that the main factors important for the observed dynamics were memory effects. We also proposed a simple algorithm to estimate the dynamics of hyperkalemia and the activation of the K\textsubscript{ATP} current using individual patient recordings. Overall, we demonstrated that taking into account the electrophysiological changes caused by hyperkalemia and hypoxia, the results of [13] can be fully explained, indicating that acidosis has only a minor effect on VF activation during the first few minutes of ischemia.
Methods

Electrophysiological model

Cardiac cellular electrophysiology was modeled using the TP06 model of human ventricular cardiomyocyte [15,17] coupled into a monodomain model for cardiac tissue. In this model the main equation determining the propagation of transmembrane voltage is given by:

\[ \frac{\partial V}{\partial t} = \nabla \cdot \left( D \nabla \left( \frac{\partial V}{\partial x} \right) \right) - I_{\text{ion}}(V, \ldots) \]  

(1)

where \( V \) is transmembrane voltage, \( D \) a diffusion tensor and \( I_{\text{ion}} \) is the sum of ionic currents:

\[ I_{\text{ion}} = I_{\text{Na}} + I_{\text{K1}} + I_{\text{Na} + I_{\text{K1}} + I_{\text{Ks}} + I_{\text{CaL}} + I_{\text{NaCa}} + I_{\text{NaK}} + I_{\text{pCa}} + I_{\text{pK}} + I_{\text{bCa}} + I_{\text{bNa}} + I_{\text{KATP}} \]  

(2)

where all currents except \( I_{\text{KATP}} \) were described by the equations from [18,15], and \( I_{\text{KATP}} \) was a new current that we introduced to take the effects of hypoxia into account. For most parameters we used the values listed in Table 1 and Table 2 of [15] corresponding to epicardial cells. For the five parameters listed in Table 2 of [15] we used the values corresponding to “Slope 1.8” for 3D simulations, and “Slope 1.1” in 2D simulations for finding the restitution curves. We used parameters corresponding to all three different slopes: 1.1, 1.4 and 1.8, for analyzing the stability of re-entry in 2D.

Ischemia was introduced to the model by changing the parameters corresponding to each of the three ischemia components described above. To model hyperkalemia we varied \( [K^+]_o \) in the range 5–10 mM. Acidosis was taken into account by decreasing the maximum conductivity of sodium and L-type calcium currents within the limits of 20–100%. The effect of hypoxia was represented by a novel description of the ATP dependent potassium current in the human heart.

Table 1 summarizes the changes to the TP06 model due to ischemia.

ATP-dependent potassium current

We developed our model of the \( I_{\text{KATP}} \) current based on the results of \textit{in vitro} experiments [16]. Figure 1 shows measured current-voltage dependencies of \( I_{\text{KATP}} \) for two different extracellular potassium concentrations. We fitted these data using functions with a power dependency on \( [K^+]_o \) concentration and exponential functions of voltage, similar to functions used in [19,20] for \( I_{\text{KATP}} \). Our expression for human \( I_{\text{KATP}} \) was:
dependency unspecified, changing the value of APD and CV restitution, a dynamic property of cardiac tissue important for the modeling framework.

For all types of the media we used ‘no flux’ boundary conditions:

$$\sum_{ij} n_i D_{ij} \frac{\partial V}{\partial x_j} = 0$$

where $n_i$ are the coordinates of a vector that is normal to the boundary.

To solve the differential equations we used a finite difference approach. For both 2D and 3D simulations we introduced a rectangular mesh of about one million points. To approximate the diffusion term we used a stencil of 17 grid points for 2D and 17 points for 3D. We used an explicit first order Euler method to solve the discretized system, which for 2D tissue was:

$$\frac{V_{ij}^{n+1} - V_{ij}^n}{\tau} = \frac{1}{h^2} \sum_{ij} w_{ij}^h V_{ij} + I_{ion}(V_{ij}, \ldots)$$

where the time step $\tau = 0.02$ ms, $h$ was the space step, and $w_{ij}^h$ weights corresponding to the diffusion tensor at location $ij$. The space step was 0.25 mm for 2D simulations, and 0.5 mm for 3D simulations [23]. The gating variables in the TP06 model were integrated using the Rush and Larsen approach [24]. To cache the results returned by functions that were only voltage dependent we used pre-computed look-up tables.

The model was implemented using the C and C++ programming languages with OpenMP extensions for parallelization. We mainly used the Intel ICC compiler toolkit. The code for 2D was run on an Intel Core i7-3930K (3.20 GHz) machine, and the 3D code for human ventricles was run on dual-processor Intel Xeon E5-2650 (2.0 GHz) machines.

Results

APD and CV restitution

In a model of a thin strip of human cardiac tissue, we studied how the different components of ischemia influence APD and CV restitution. A model of human ventricles presented in [21]. This model takes anisotropy into account by reconstructing the fiber direction field described in [22] and assuming that the diffusion coefficient across the fiber $D_f$ is 4 times less than the diffusion coefficient along fibers $D_t$, which was set to 1.54 cm$^2$/s. For 3D simulations the components of the diffusion tensor were given by:

$$D_{ij} = (D_t - D_f)\tau_i \tau_j + D_f \delta_{ij}$$

where $\tau_i$ are the coordinates of a normalized vector oriented along the fibers.

Numerical methods

For 1D and 2D simulations we solved equation (1) for homogeneous and isotropic tissue with the diffusion tensor $D_\delta = D_\delta \delta_\delta$ where $D_\delta$ was taken to be 1.54 cm$^2$/s and $\delta_\delta$ is the Kronecker delta. For these parameter values, the velocity of propagating plane waves at a stimulation frequency of 1 Hz was 72 cm/s.

For 3D whole heart simulations we used an anatomically based model of human ventricles presented in [21]. This model takes anisotropy into account by reconstructing the fiber direction field described in [22] and assuming that the diffusion coefficient across the fibers $D_f$ is 4 times less than the diffusion coefficient along fibers $D_t$, which was set to 1.54 cm$^2$/s. For 3D simulations the components of the diffusion tensor were given by:

$$D_{ij} = (D_t - D_f)\tau_i \tau_j + D_f \delta_{ij}$$

where $\tau_i$ are the coordinates of a normalized vector oriented along the fibers.
onset of re-entry and for the stability of re-entrant waves [25,26]. We followed the same approach as in [14], but with a different formulation of $I_{KATP}$.

Restitution and dispersion curves were obtained using an S1S2 protocol, in a $10 \times 2.5$ cm sheet of 2D simulated tissue. Super-threshold stimuli were delivered along one short edge of the sheet, and measurements of APD were made at a distance of 2.5 cm from the stimulated edge. The basic cycle length (BCL) for S1 stimuli was 1000 ms. The ten S1 stimuli were followed by a single S2 stimulus. The duration between the last S1 and S2 was decremented from 1000 ms until there was no response to the S2 stimulus at the point of measurement. The restitution curves are shown in Figure 2.

In Figure 2A the effect of hyperkalemia on APD restitution is shown, indicating that APD decreases with increasing $[K^+]_o$. This effect is due to $I_{K1}$ and $I_{K2}$ currents, which depend directly on $[K^+]_o$ and become larger once $[K^+]_o$ is increased, accelerating the repolarization process. The effect of hyperkalemia on conduction velocity is shown in Figure 2D. We see that the conduction velocity decreases with increasing $[K^+]_o$. This is because an increase in $[K^+]_o$ shifts the resting potential to more positive values suppressing the sodium current and, consequently, reducing the excitability of the cell.

Figures 2A and 2D also show how the minimal APD—APD$_{\text{min}}$—value (the left most point of the restitution curve) depends on $[K^+]_o$. APD$_{\text{min}}$ increases substantially with hyperkalemia. This fact can be important as APD$_{\text{min}}$ is considered as one of the main factors determining the complexity of pattern of excitation during fibrillation [21,27].

Figures 2B and 2E illustrate the effect of acidosis (modeled as reduction of $I_{Na}$ and $I_{CaL}$) on APD and CV restitution. Both APD and CV decrease with acidosis, because acidosis diminishes the depolarizing currents available to the cell. However, the value of the APD$_{\text{min}}$ showed almost no dependency on acidosis, unless the conductance of the $I_{Na}$ and $I_{CaL}$ channels was reduced to 20% of their default values.

Figures 2C and 2F show the effect of hypoxia, modeled as activation of $I_{KATP}$. Both APD and APD$_{\text{min}}$ become substantially shorter since hypoxia activates a strong depolarizing current. The value of CV showed almost no dependency on hypoxia because hypoxia does not affect $I_{Na}$.

Figure 2 shows that each component of ischemia acted to reduce the slope of restitution. Steep restitution is considered as one of the main mechanisms contributing to the breakup of re-entry in VF [25,26]. According to the restitution hypothesis, flattening restitution prevents possible spiral wave breaks from happening, and therefore prevents formation of new arrhythmia sources.

**Effect of ischemia on arrhythmia sources in 2D**

We studied how the different components of ischemia influence the dynamics and stability of re-entrant waves in 2D—spiral waves. In these simulations we varied the slope of the restitution curve for non ischemic conditions by changing five parameters of

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**Figure 2. A, B, C: APD restitution curves for hyperkalemia, acidosis and hypoxia respectively. D, E, F: CV dispersion relations for hyperkalemia, acidosis and hypoxia respectively.** P denotes the sum of diastolic interval (DI) and APD at the point of measurement for the pulse generated by the S2 stimulus.

doi:10.1371/journal.pcbi.1003891.g002
our model, as described in the Methods. We used three different sets of parameters, which correspond to a slope of 1.1, 1.4 and 1.8 under normal conditions. For initiation of a spiral we used an S1S2 protocol: at first, the S1 stimulus was delivered along one side of the medium, then the S2 stimulus was applied after the wave had passed half of the medium.

Figure 3 shows the resulting patterns of transmembrane voltage after 10 s of simulated activity for each parameter set and under each component of ischemia. For tissue with a restitution slope of 1.1 under normal conditions, the spiral remained stable regardless of ischemia. For a restitution slope of 1.4 (the second row), a complex fibrillatory pattern developed under normal conditions resulting from breakup of the initial spiral wave, however each ischemia factor prevented breakup. The third row gives an example for a relatively high restitution slope of 1.8 under normal conditions. In this case the spiral wave broke into fibrillation under both normal and ischemic conditions. However, it was still possible to stabilize the spiral rotation by further increasing the ischemia parameters. The bottom row in Figure 3 corresponds to that case: as ischemia became more severe the activation pattern did not evolve into fibrillation.

Overall, all three components of ischemia acted to stabilize re-entry, and to prevent the breakdowns that can happen under normal conditions. This is in line with the prediction that we derived from analyzing the restitution curves in the previous section, which was based on flattening of the restitution curves due to ischemia.

Our next step was to understand the change of the main dynamical characteristics of cardiac tissue with ischemia and to compare these with the activation patterns observed in the human heart [13]. Therefore, we performed simulations using an

| Restitution slope for normal conditions | Normal conditions | Hyperkalemia $[K^+]_o = 6$ mM | Acidosis $G_{Na,CaL} = 80\% \cdot G^0_{Na,CaL}$ | Hypoxia $f_{ATP} = 0.1\%$
|---|---|---|---|---|
| 1.1 | | | | |
| 1.4 | | | | |
| 1.8 | | | | |

| Restitution slope for normal conditions | Normal conditions | Hyperkalemia $[K^+]_o = 7$ mM | Acidosis $G_{Na,CaL} = 60\% \cdot G^0_{Na,CaL}$ | Hypoxia $f_{ATP} = 0.3\%$
|---|---|---|---|---|
| 1.8 | | | | |

**Figure 3. Stability of spirals under ischemic conditions.** Greyscale shows membrane voltage, with light grey indicating depolarized tissue. doi:10.1371/journal.pcbi.1003891.g003
anatomically detailed model of the human ventricles developed in [21]. To explain our results we shall use information on 1D and 2D wave propagation collected in the previous sections.

**VF in an anatomical model of the human ventricles incorporating simulated global ischemia**

**Wave patterns.** The three components of ischemia were modeled as described in Table 1. VF was induced using an S1S2 protocol which resulted in a subsequent spiral wave breakup and onset of the fibrillatory pattern as described in [21]. It was shown that this protocol reproduces the main features of VF in the human heart [21,10]. The resulting state after 5 s of initialization was saved and used as an initial condition for subsequent simulations to investigate the effect of ischemic components on VF dynamics.

Figure 4 shows example snapshots of activation in the anatomically detailed model for both normal and ischemic conditions, with each ischemic component set to values corresponding to those obtained from the human recordings, as we will show later.

Figures 4A–D show typical excitation patterns during simulated VF, both in terms of wavefronts (A: front view, B: lateral view) and activated regions for which the voltage is higher than APD₉₀ level (C, D). Here we have a complex fibrillatory pattern characterized by multiple re-entrant waves. For every subsequent simulation, we started from this initial state and gradually (within 200–800 ms) changed the corresponding parameters from their normal values to the values corresponding to the different ischemic conditions, and then computed for a further 15 s of model time. Examples of fibrillation patterns we obtained are shown in Figure 4E–P.

Figures 4E–H give the resulting picture for hyperkalemia with \([K^+]_o = 7\) mM. The complexity of activation was slightly lower in comparison with the initial state, with fewer wavefronts. Figures 4G–H show the amount of activated tissue for this snapshot. The ratio of green to red was larger for hyperkalemia, which indicates that the tissue became less excitable. Similar results can be seen for hypoxia \(f_{\text{ATP}} = 0.1\%\) in Figures 4M–P. The medium was less excitable and the complexity of VF was slightly lower compared to normal.

Acidosis had a less pronounced effect on both excitability and pattern complexity. Figures 4I–L give an example of fibrillation under acidosis conditions where the conductivity of \(I_{\text{Na}}\) and \(I_{\text{CaL}}\) were reduced to 80%.

Period of VF. The period of VF in each simulation was determined using a discrete Fourier transform of time series of transmembrane voltage obtained from 10000 locations in the heart model for the last 7 s of model time, and obtaining the period from the frequency of the largest peak in the spectrum. We found that the period of fibrillation showed almost no dependence on acidosis. Even for a severe drop of \(I_{\text{Na}}\) and \(I_{\text{CaL}}\)
The dependence of the average period of VF on hypoxia and hyperkalemia is shown in Figure 5. The period of fibrillation increased under hyperkalemia, and decreased under hypoxia. The effect of hyperkalemia was more pronounced for smaller values of \( f_{ATP} \). Also, a saturation effect can be seen: for larger values of \( f_{ATP} \) the effect of additional hypoxia was relatively small.

At the first glance, our observation that hyperkalemia acted to increase the period of VF contradicts the results of our 1D studies, which showed that APD decreased with increase of \( [K^+]_o \). However, as in the case of acidosis, the most important factor here is \( \text{APD}_{min} \) which increases with hyperkalemia (Figure 2A) and, therefore, explains the observed increase in the period of VF.

Explanation of the observed decrease of period under hypoxia is straightforward. The main effect of hypoxia is shortening of the action potential. Indeed, we see in Figure 2C that \( \text{APD}_{min} \) becomes smaller under hypoxia. This shortening of APD results in a decrease of the period of VF. The effect of hyperkalemia turned out to be more pronounced for smaller values of \( f_{ATP} \).

We used the data of Figure 5 to estimate the effects of these two factors of ischemia on the period change during ischemia observed in [13].

**Estimation of the relative contribution of hyperkalaemia and hypoxia to patient data**

Our results allowed us to estimate the extent of hyperkalaemia and hypoxia for each patient involved in the clinical study [13]. We used the dependency of how DF (1/period) changes throughout experimental VF for individual patients. One of these recordings is shown in Figure 6. The red region corresponds to normal perfusion of the heart, blue to global ischemia and the green to reperfusion. As one can see, DF gradually decreases during ischemia and then increases abruptly after reperfusion to an even higher level that it was initially.

For each patient we used three points on this graph: DF at the beginning of ischemia (30 s, point A), DF at the end of ischemia (180 s, point B) and DF at the end of the experiment (210 s, point C). Ischemia takes place in between points A and B, thus \( [K^+]_o \) and \( f_{ATP} \) are increasing there. At point B we expect to have the highest values for these quantities.

At point B reperfusion starts, resulting in a rapid elevation of DF. Figure 5 indicates that hypoxia acts to shorten the VF period, thus it is likely to be responsible for this change. Therefore we assumed that the recovery of hypoxic \( K_{ATP} \) channels during reperfusion is slower than the recovery of extracellular potassium concentration. We assumed that \( [K^+]_o \) had returned to a normal value by the point C whereas \( f_{ATP} \) remains at the same level as at point B. Thus, knowing DF at point C, we can find the fraction of open \( K_{ATP} \) channels at that moment. Using this value and the value of DF we can estimate \( [K^+]_o \) at point B. The results we obtained using this algorithm are given in first three columns of Table 2. The results shown in columns 4–7 of the table deal with a possible partial recovery of \( f_{ATP} \) and will be described in the discussion section. The results in the last two columns are based on the assumption that \( K_{ATP} \) channels open even before the beginning of ischemia. This will also be described in the discussion section.

We see that the estimated \( [K^+]_o \) concentrations for different patients vary in the range from 6.5 to 8.0 mM, while the fraction of activated \( K_{ATP} \) channels \( f_{ATP} \) does not go beyond 0.1%.

We also estimated how hypoxia and hyperkalaemia change in the course of ischemia. This problem of fitting does not have a unique solution because the period of fibrillation depends on two parameters, while we have only one period dependency for each patient. To account for that we assumed that the dependencies of \( [K^+]_o \) and \( f_{ATP} \) are monotonic with time, and these values can never decrease during ischemia. Then we wrote the relation between the rate of change of these values:

\[
\frac{dT}{dt} = \frac{\hat{c}T}{\hat{c}[K^+]_o} \frac{d[K^+]_o}{dt} + \frac{\hat{c}T}{\hat{c}f_{ATP}} \frac{df_{ATP}}{dt},
\]

where \( T(t) \) is the period of fibrillation over time (from the clinical results) and \( T([K^+]_o, f_{ATP}) \) is the dependency of VF period on the...
ischemia components, obtained from Figure 5. Our goal was to determine the patient specific functions $\frac{[K^+]_o(t)}{[C^138o]}$, based on known $T(t)$, $T([K^+]_o,f_{\text{ATP}})$ and boundary conditions from Table 2. To solve (5) we needed to impose an additional constraint on our functions. Thus we assume that

$$\frac{d[K^+]_o}{df_{\text{ATP}}} = \frac{[K^+]_o,B - [K^+]_o}{f_{\text{ATP}},B - f_{\text{ATP}}}$$

(6)

where $[K^+]_o,B$ and $f_{\text{ATP},B}$ are the values we fitted for point B. This constraint ensured the values of hypoxia and hyperkalemia tended to reach those values at point B. The results we obtained using this approach for two patients are shown in Figure 7. The rest of the results for all 10 patients are available in Figures S1–S10. We see that we can fit the clinical data with smooth monotonic functions using our approach. However, as our constraint (6) cannot be justified from biological background, these fits can be considered as a conjecture rather than established results.

**Number of re-entrant sources**

The study of [13] describes how the number of epicardial phase singularities and the number of wave fronts change during the course of ischemia. These values have approximately the same dynamics and to understand the results we studied the change in the number of phase singularities. Phase singularities are points where filaments intersect the surface of the heart [7,10]. As was shown in [10], the number of filaments obtained from simulations in an anatomically detailed model of human ventricles provide a good estimate of the number of phase singularities on the surface of the heart. We counted the number of filaments in our model in normal conditions and under different factors of ischemia.

Figure 8 shows four examples of how the number of filaments depends on time under conditions that we expect to correspond to those observed for the first 2.5 minutes of ischemia. Figure 8A shows filament numbers during normal conditions. We see dynamics similar to that reported in [21]. Figures 8B and C correspond to hyperkalemia $[K^+]_o \approx 7$ mM and hypoxia $f_{\text{ATP}} \approx 0.1\%$, respectively. We see that compared to normal condition there is a tendency for decreasing of the number of filaments. Finally, Figure 8D gives an example where simulated VF terminated spontaneously, although the number of filaments stayed relatively high until immediately prior to termination. For that simulation we used $[K^+]_o \approx 8$ mM and $f_{\text{ATP}} = 0.4\%$.

The mean number of filaments in the two parameter space—hyperkalemia and hypoxia—for a wider range of parameters is given in Figure 9. The error bars in this figure correspond to the standard deviations of root mean square. The point with $[K^+]_o = 5.4$ mM and $f_{\text{ATP}} = 0$ corresponds to normal conditions. We can see that for values of hypoxia $f_{\text{ATP}} < 0.1\%$ the complexity of the pattern slightly decreases. However, it increases for larger values of hypoxia. All these results differ substantially from our observation from the 2D patterns (see Figure 3). In 2D, we observed that every single component of ischemia suppressed breakup, which would correspond to a decrease of filament numbers and termination of fibrillation in our whole heart simulations. However, this figure shows that for 3D an increase in fibrillation complexity as the degree of ischemia increases.
This is a surprising result that can be explained in the following way. The 2D simulations always started from a single spiral, and breakup was self induced. In 3D, however, we started from an initial pattern with multiple re-entrant waves. This complex pattern was able to maintain itself in spite of the presence of simulated ischemia. To confirm this observation, we performed simulations in 2D, starting from a developed VF pattern. These simulations showed that even in 2D, the complex pattern can persist compared with initiation with the S1S2 protocol of Figure 3.

The results of the clinical research are presented in Table 3. The minimal and maximal value of the number of phase singularities on the surface of the heart are given there for both the beginning and the end of ischemia. As shown, there are large deviations in PS numbers making it difficult for us to give conclusive statements from these data. However, they do indicate relative conservation on PS numbers during ischemia.

If we now compare these results with results on filament numbers shown in Figure 9 we can conclude the following. In the range of ischemic components which we estimated for the individual patients (\(\frac{1}{2}C_138\) mM, \(f_{ATP} < 0.1\%\)) we see some tendency for a decrease in the filament numbers in our simulations. However, this decrease is minimal and to some extent can be considered as conservation of filament numbers in a

![Figure 7. Two examples of how hypoxia and hyperkalemia evolve during course of ischemia.](doi:10.1371/journal.pcbi.1003891.g007)

![Figure 8. Change of the number of filaments in course of time under different ischemia factors obtained from numerical simulations.](A: normal conditions, B: hyperkalemia, C: hypoxia, D: example of spontaneous fibrillation stop. doi:10.1371/journal.pcbi.1003891.g008)
way similar to that we see in clinical data. However, our simulations indicate that we can expect an increase in filament numbers if ischemia is more prolonged or if $I_{KATP}$ channels open to a greater extent.

**Discussion**

The aim of this study was to use a model of cardiac electrical activity to gain insight into the mechanism underlying the organization of ventricular fibrillation observed in the in-situ human heart with global cardiac ischemia. We performed numerical simulations using a detailed electrophysiological model of cardiac cells and tissue under normal and ischemic conditions. This model was examined in pseudo-1D and 2D tissue geometries, and in an anatomical model of human ventricles which takes the geometry and anisotropy of cardiac tissue into account. As in [5,20,19] we subdivided the effects of ischemia into three main syndromes: hyperkalemia, acidosis and hypoxia. To represent the activation of $K_{ATP}$ channels in hypoxia, we developed a new model of the human $K_{ATP}$ channel, based on recordings of [16].

Our 1D and 2D studies demonstrated that each component of ischemia results in a shortening of APD under the same base cycle length, which is in accordance with the results of other modeling research [19]. Our results for CV restitution (dispersion relation) are similar to those obtained in [20] with a small difference: the CV does not depend monotonically on hyperkalemia. As can be seen in Figure 2D, in our simulations CV increases with hyperkalemia for lesser values of $\frac{[K^+]_o}{C_{138}}$ whereas for higher ones it decreases. This supernormal conductance effect was not observed in studies of [20], which may be due to a different stimulation protocol. Overall, the restitution curves we obtained are very similar to [14].

According to the restitution hypothesis, our observation that the steepness of APD restitution curves decreases with ischemia makes dynamical instabilities less likely to occur, and wavebreak leading to multiple wavelet VF less probable [26]. We confirmed this in our 2D simulations. In particular, we found that it was sufficient to increase any of the ischemic factors within their physiological range to prevent break up that can occur under non ischemic conditions. This is also in line with the simulations on effects of activation pattern due to restitution steepness by [26,28].

The main part of the present study deals with simulations in a 3D anatomically accurate model of the ventricles of the human heart for situations that resemble the clinical set up used in [13]. Our aim was to describe how the organization of simulated VF is altered due to cardiac ischemia. We found that acidosis had almost no effect on the activation rate, while hyperkalemia decreased and hypoxia increased the frequency of VF. Studies of this effect were not previously performed in anatomical models, although they are similar to 2D simulations and experiments in animal models. The effect of hyperkalemia on the period of spiral wave rotation was studied by [29] who obtained similar dependency but only for concentrations of $[K^+]_o$ larger than 8 mM. The same effect of hypoxia was investigated by [30] on canine ventricular slices and later by [31] on swine hearts. The results obtained in their experiments are in accordance with our simulations.

The dependence of activation rate on ischemic factors could be predicted by looking at our APD restitution curves. Fibrillatory

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**Table 3.** Minimal and maximal values of the number of phase singularities during the first and the last ten seconds of ischemia for the patients studied in [13].

| Patient | Before ischemia | After ischemia |
|---------|-----------------|----------------|
| H055    | 2–17            | 1–18           |
| H057    | 2–15            | 2–20           |
| H058    | 2–22            | 3–26           |
| H059    | 2–22            | 1–18           |
| H060    | 4–22            | 5–24           |
| H062    | 1–14            | 1–19           |
| H063    | 1–13            | 2–15           |
| H065    | 1–11            | 1–18           |
| H066    | 2–18            | 2–22           |

doi:10.1371/journal.pcbi.1003891.t003

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**Figure 9.** The mean number of filaments of a fibrillation pattern averaged over time of simulation after applying ischemic conditions. The error bars indicate the standard deviation of root mean square. doi:10.1371/journal.pcbi.1003891.g009
excitation patterns have a very high activation rate, and thus the
leftmost regions of restitution curves are of considerable impor-
tance since they correspond to high activation rate and short
diastolic interval. If we look, for instance, at Figure 2A we see that
the leftmost points shift to the right with hyperkalemia. Thus the
activation frequency of fibrillation decreases. The minimal possible
values of APD, which are given by the left most points, are called
APD_{min}. Thus this finding once again emphasizes a particular
importance of these APD_{min} on VF organization which has also
been demonstrated in other studies [23,27]. Note, that the
APD_{min} point is achieved at the minimal possible period of
stimulation, which is, perhaps, more directly related to the period
of excitation during VF.

With our 3D simulations we were able to construct a model of
the dependency of VF period on both hyperkalemia and hypoxia.
This model, along with additional assumptions about the relative
rates at which the different components of ischemia develop
allowed us to propose a way to estimate the extracellular potassium
concentrations and the fraction of open KATP channels for the
recordings in the clinical study [13]. The estimated external
potassium concentrations correspond to experimental data show-
ing a rate of $[K^+]_o$ change of about 0.5–1 mM per minute [2].

The most important assumption in this study was that the rate
of recovery from hypoxia during reperfusion is slower than the rate
of recovery from hyperkalemia. Experimental data from porcine
hearts [32] show that the complete recovery of $[K^+]_o$ during
reperfusion can occur within less than one minute. Unfortunately
the data available on the recovery of KATP channels are controver-
sial which makes it difficult to assess whether our assumption is correct. Data
from the the human heart [13,33] obtained at the whole organ
level show that, even after 5 minutes of reflow, less than 70% of
the APD shortening (which was assumed to be an effect of KATP
opening) had been reversed. On the other hand, experimental
data from single cells demonstrate that recovery can be very fast,
of the order of a few seconds [34,35].

To investigate an effect of this possible partial recovery of
hypoxia we estimated how our estimates change if we assume that
after 30 seconds of reperfusion either 25% or 50% of KATP
had been recovered. These estimates are shown in the columns 4–7 of
Table 2. The second and the third columns are given for the case
of no recovery. As we can see, partial recovery does not have a
pronounced effect on the estimated values of hyperkalemia during
ischemia. Indeed, for most of the patients the error in $[K^+]_o$
estimation due to this partial recovery is about 0.3–0.4 mM.

We have also proposed a method of fitting the change in $[K^+]_o$
and $f_{ATP}$ for each individual patient during global cardiac
ischemia. Note, however, this problem is nontrivial because its
exact solution requires more experimental details on the speed of
various processes during ischemia. The approach that we followed
is based on solving a boundary problem for an ordinary
differential equation (ODE), expanded with one artificial
ODE that provided a relation between relative change of the
hyperkalemia and hypoxia factors. We also tried other, less sophisticated
methods to target this problem, such as linearly varying one of the
factors and finding the time course for the other one. The
deficiency of these simple methods was that they violated the
assumption that ischemia factors grow monotonically for some of
the data. Therefore we followed our approach, although it is not
based on any mechanistic background. As soon as more data on
the effects of ischemia in the human heart are available, our
approach could be improved by adding more constraints on the
solution.

Our clinical data show that in all recordings there was an initial
increase of the dominant frequency that happens during the first
30 s of the study under normal perfusion. This effect was also
obtained in other studies [9,36]. A possible explanation for this
effect is the following. Since ventricular fibrillation is an energy
consuming process, it may shift the [ADP] to [ATP] ratio, and
hence diminish the energy capabilities of the cells. This would lead
to opening of KATP channels, and result in an increased DF. To
examine this possible explanation, we modified our fitting method,
taking the DF at the beginning of the recording as corresponding
to normal conditions. The maximum values for hypoxia and
hyperkalemia for this case are shown in the last the columns 8–9 of
Table 2. We see that this alternative assumption leads to an
increased number of activated of KATP channels, whereas the
extracellular potassium concentration remains relatively un-
changed. We also adapted this assumption for our fitting algorithm
describing how hypoxia and hyperkalemia evolve in the course of
ischemia. The results for two patients are given in Figure 10.

Finally, we studied the effects of global cardiac ischemia on the
complexity of activation patterns during simulated VF. For that we
used an established VF activation pattern as an initial condition for
the simulations. This type of initial condition differs from most of
the previous studies of the restitution based fibrillation, where a
single spiral is used for the initial condition and the conditions
under which this spiral develops into fibrillation are in question.
We found that using established VF as an initial condition showed
that flattening of the restitution curve caused by ischemia does not
usually lead to VF termination, but rather results in a small
number of stable re-entrant sources. We think that this finding
does not contradict to the results obtained on spiral breakups in
2D. Indeed, flattening of restitution curves just prevents the onset
of VF, however, this process is not directly related to the
elimination of multiple existing spirals. In fact, flattening of
the restitution curves decreases dynamical instabilities and makes
the rotation of spirals more stable, thus it is even more likely to
stabilize multiple spirals in the tissue. The tendency for a
fibrillation pattern to become more organized due to global
ischemia was first demonstrated experimentally by [37] on rabbit
hearts. In addition, a recent experimental study on the effects of
hypoxia in swine hearts by [31] has also shown that activation of
KATP channels makes the fibrillation pattern more stable and stops
rotors from meandering. However, a study by [26], performed also
on swine hearts, argues that flattening of the restitution curve does
favour VF termination, though, to achieve that flattening they
used a potassium channel antagonist—breylinium—which has an
effect opposite to hypoxia.

Limitations
There are several factors that were not taken into account in this
study. We used a homogeneous model of the heart which does not
take into account the differences between epicardial, endocardial
and M-cells. However, as it is shown in [38] the differences
between these cell types decrease substantially at the high
frequency of excitation typical for VF patterns. Note that the
differences between different cell types are not that pronounced in
tissue due to the electrotonic effect. Next, we used a monodomain
model, to represent the cardiac tissue. Bidomain models are of
considerable importance when describing defibrillation phenom-
ena but in the absence of applied external currents the difference
between these two types is extremely small [39].

Due to numerical limitations, we were not able to reproduce full
3.5 minutes of the clinical study in our model and limited ourselves
by only 20 seconds. We did not check if the effects we studied are
also presented in other detailed electrophysiological models of
cardiomyocyte, such as [40] and [41], nor did we take into account
effects due to mechanical activity of the heart during VF.
We did not consider the change in the gap junction conductance caused by ischemia. There are several experimental studies on animal models that demonstrate that the effect on gap junction uncoupling is relatively small in first minutes of ischemia. The investigations in a rabbit papillary muscle [42] show that under ischemic conditions the intracellular resistance stays constant within the first 10–15 minutes, whereas the extracellular resistance slightly changes immediately following the onset of ischemia. However this change was not associated with a significant decrease of the conduction velocity within the first 4 minutes of ischemia. Another result obtained in isolated rat hearts [43] indicates that the absolute value of the impedance of the tissue changes for about 3 Ohm cm/min with the initial value of about 150 Ohm cm. Finally, another set of experiments on rabbit papillary muscle [44] also shows that the tissue resistance stays remarkably stable for the first 10 minutes of acute ischemia. These data allow us to assume that the process of gap junction uncoupling does not have a significant effect on the conduction velocity for the time scale of 2.5 minutes of ischemia.

We did not consider either the heterogeneity of the tissue response to ischemia nor heterogeneity in ionic channels distributions. As shown in other studies [45], heterogeneities in \( I_{\text{KATP}} \) distribution can lead to a transmural gradient of the DF in myocardium. There are several studies that show that this gradient occurs in experimental models [11,12,46]. Furthermore, global ischemia may lead to different depths of ischemia in different regions of the heart, since the endocardium is in contact with a large volume of oxygenated blood in the ventricular cavities, at least in the early stages of VF. We expect that these heterogeneities can lead to a different fibrillation mechanism (so-called mother rotor fibrillation) as well as increase in the number of filaments [47,48]. This factor may also contribute to increasing the number of phase singularities localized at the surface of the heart.

The role of Purkinje system was not considered in this study. There are several experimental studies in dog hearts [46,49], which show that the Purkinje system can play an important role in early termination of VF because it acts to increase the transmural activation rate gradient. However, a computer modeling study [45] shows that this effect is not that prominent in comparison with the effect of heterogeneity in \( I_{\text{KATP}} \) distribution.

Finally, we omitted several other effects that may occur during ischemia. In our case, depletion of ATP results only in activation of the \( K_{\text{ATP}} \) channel. We did not consider the effect of ATP on other important processes, such as functioning of the \( K_{\text{Na}} \) pump, the processes occurring at the mitochondria membranes and their effect on calcium dynamics. We did not take into account any changes of cell size or extracellular medium size occurring as a result of osmosis, and possible changes of the ionic concentrations due to it. It would be valuable to extend existing cell models to incorporate a more detailed description of ischemia, and to study the effects of ATP depletion using such a model.

Supporting Information

Figure S1  Evolution of hypoxia and hyperkalemia in the course of global ischemia for the patient H053 in [13], estimated by our fitting method. Pink: clinical data on the dominant frequency, averaged over 1 s window with standard deviation. Blue: smoothed curve that we used for our fitting. Red: extracellular potassium concentration. Green: fraction of activated \( K_{\text{ATP}} \) channels.

Figure S2  Evolution of hypoxia and hyperkalemia in the course of global ischemia for the patient H057 in [13], estimated by our fitting method. Pink: clinical data on the dominant frequency, averaged over 1 s window with standard deviation. Blue: smoothed curve that we used for our fitting. Red: extracellular potassium concentration. Green: fraction of activated \( K_{\text{ATP}} \) channels.

Figure S3  Evolution of hypoxia and hyperkalemia in the course of global ischemia for the patient H058 in [13], estimated by our fitting method. Pink: clinical data on the dominant frequency, averaged over 1 s window with standard deviation. Blue: smoothed curve that we used for our fitting. Red: extracellular potassium concentration. Green: fraction of activated \( K_{\text{ATP}} \) channels.

Figure S4  Evolution of hypoxia and hyperkalemia in the course of global ischemia for the patient H059 in [13], estimated by our fitting method. Pink: clinical data on the dominant frequency, averaged over 1 s window with standard deviation. Blue: smoothed curve that we used for our fitting. Red: extracellular potassium concentration. Green: fraction of activated \( K_{\text{ATP}} \) channels.
**Figure S5** Evolution of hypoxia and hyperkalemia in the course of global ischemia for the patient H060 in [13], estimated by our fitting method. Pink: clinical data on the dominant frequency, averaged over 1 s window with standard deviation. Blue: smoothed curve that we used for our fitting. Red: extracellular potassium concentration. Green: fraction of activated $K_{ATP}$ channels.

**Figure S6** Evolution of hypoxia and hyperkalemia in the course of global ischemia for the patient H062 in [13], estimated by our fitting method. Pink: clinical data on the dominant frequency, averaged over 1 s window with standard deviation. Blue: smoothed curve that we used for our fitting. Red: extracellular potassium concentration. Green: fraction of activated $K_{ATP}$ channels.

**Figure S7** Evolution of hypoxia and hyperkalemia in the course of global ischemia for the patient H063 in [13], estimated by our fitting method. Pink: clinical data on the dominant frequency, averaged over 1 s window with standard deviation. Blue: smoothed curve that we used for our fitting. Red: extracellular potassium concentration. Green: fraction of activated $K_{ATP}$ channels.

**Figure S8** Evolution of hypoxia and hyperkalemia in the course of global ischemia for the patient H064 in [13], estimated by our fitting method. Pink: clinical data on the dominant frequency, averaged over 1 s window with standard deviation. Blue: smoothed curve that we used for our fitting. Red: extracellular potassium concentration. Green: fraction of activated $K_{ATP}$ channels.

**Figure S9** Evolution of hypoxia and hyperkalemia in the course of global ischemia for the patient H065 in [13], estimated by our fitting method. Pink: clinical data on the dominant frequency, averaged over 1 s window with standard deviation. Blue: smoothed curve that we used for our fitting. Red: extracellular potassium concentration. Green: fraction of activated $K_{ATP}$ channels.

**Figure S10** Evolution of hypoxia and hyperkalemia in the course of global ischemia for the patient H066 in [13], estimated by our fitting method. Pink: clinical data on the dominant frequency, averaged over 1 s window with standard deviation. Blue: smoothed curve that we used for our fitting. Red: extracellular potassium concentration. Green: fraction of activated $K_{ATP}$ channels.

**Author Contributions**
Conceived and designed the experiments: IVK AVP. Performed the experiments: IVK MPH PT AVF. Analyzed the data: IVK RHC AVF. Contributed reagents/materials/analysis tools: IVK AVF. Wrote the paper: IVK RHC MPH CPB DJP PT AVF.

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