Leak current, even with gigaohm seals, can cause misinterpretation of stem cell-derived cardiomyocyte action potential recordings

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Aims
Human-induced pluripotent stem cell-derived cardiomyocytes (iPSC-CMs) have become an essential tool to study arrhythmia mechanisms. Much of the foundational work on these cells, as well as the computational models built from the resultant data, has overlooked the contribution of seal–leak current on the immature and heterogeneous phenotype that has come to define these cells. The aim of this study is to understand the effect of seal–leak current on recordings of action potential (AP) morphology.

Methods and results
Action potentials were recorded in human iPSC-CMs using patch clamp and simulated using previously published mathematical models. Our in silico and in vitro studies demonstrate how seal–leak current depolarizes APs, substantially affecting their morphology, even with seal resistances ($R_{seal}$) above 1 GΩ. We show that compensation of this leak current is difficult due to challenges with obtaining accurate measures of $R_{seal}$ during an experiment. Using simulation, we show that $R_{seal}$ measures (i) change during an experiment, invalidating the use of pre-rupture values, and (ii) are polluted by the presence of transmembrane currents at every voltage. Finally, we posit that the background sodium current in baseline iPSC-CM models imitates the effects of seal–leak current and is increased to a level that masks the effects of seal–leak current on iPSC-CMs.

Conclusion
Based on these findings, we make recommendations to improve iPSC-CM AP data acquisition, interpretation, and model-building. Taking these recommendations into account will improve our understanding of iPSC-CM physiology and the descriptive ability of models built from such data.

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This is an Open Access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.
Human-induced pluripotent stem cell-derived cardiomyocytes (iPSC-CMs) are an emerging tool in the study of cardiac arrhythmia mechanisms. Their immature and heterogeneous action potential phenotype complicates the interpretation of experimental data and has slowed their acceptance in industry and academia. We suggest that the leak current caused by imperfect pipette membrane seal during single-cell patch clamp experiments is partly responsible for causing this heterogeneity and the appearance of immaturity.

Using in vitro experiments and computational modelling, we show that this seal–leak current affects iPSC-CM action potential morphology, even under ‘ideal’ experimental conditions.

Based on these findings, we make recommendations that should be considered when interpreting, analysing and fitting iPSC-CM data.

**What’s new?**

- Human-induced pluripotent stem cell-derived cardiomyocytes (iPSC-CMs) are an emerging tool in the study of cardiac arrhythmia mechanisms.
- Their immature and heterogeneous action potential phenotype complicates the interpretation of experimental data and has slowed their acceptance in industry and academia.
- We suggest that the leak current caused by imperfect pipette membrane seal during single-cell patch clamp experiments is partly responsible for causing this heterogeneity and the appearance of immaturity.
- Using in vitro experiments and computational modelling, we show that this seal–leak current affects iPSC-CM action potential morphology, even under ‘ideal’ experimental conditions.
- Based on these findings, we make recommendations that should be considered when interpreting, analysing and fitting iPSC-CM data.

**Introduction**

Human-induced pluripotent stem cell-derived cardiomyocytes (iPSC-CMs) are a renewable and cost-effective model for studying genetic disease mechanisms, drug cardiotoxicity, and inter-patient variability. Computational approaches have been developed to translate experimental results from iPSC-CMs to make predictions in adult cardiomyocytes. Such work attempts to bridge the critical gap that remains between the physiology of iPSC-CMs and excised adult human cardiac cells.

Whilst iPSC-CMs have transformed many areas of cardiac arrhythmia research, phenotypic heterogeneity and immaturity continue to stymie their potential impact. Investigating sources of these limitations and their biological implications is important as iPSC-CMs (and mechanistic models describing their behaviour) are used to inform increasingly complex clinical decisions. Studies of iPSC-CMs in a single-cell patch clamp context have indicated that their depolarized, highly varying resting membrane potential is primarily due to decreased inward rectifier potassium current (Ikr) and increased funny current (If) compared with adult cardiomyocytes.

Recently, findings from Horváth et al. and Van de Sande et al. indicate that the heterogeneous and depolarized resting membrane potential is also due, far more than previously thought, to a simple seal–leak current (Ileak). Relative to electrically coupled iPSC-CMs, they show a substantial depolarization in the resting membrane potential in isolated iPSC-CMs despite some cells having similar If densities to human adult cardiomyocytes. These findings indicate that Ileak plays an important role in iPSC-CM AP morphology during single-cell patch clamp experiments.

Ileak is inversely proportional to the seal resistance (Rseal) formed between the micropipette tip and cell membrane during patch clamp experiments. A sufficiently large Rseal is expected to limit Ileak’s effect on AP morphology. Upon reviewing single-cell electrophysiological iPSC-CM studies, including those used to build iPSC-CM computational models, we found that studies do not report either an Rseal acceptance criteria or an average Rseal < 3 GΩ.

In this study, through *in vitro* experiments and computational modelling, we show that Ileak affects iPSC-CM AP morphology, even above the Rseal values usually deemed acceptable in the literature. We show that Rseal cannot be easily compensated because it cannot be accurately measured during an experiment. Additionally, we posit that the background sodium current (IhNa) in iPSC-CM models may be overestimated and mimic the effects of leak on AP morphology. Ultimately, we argue that leak current should be considered when interpreting, analysing, and fitting iPSC-CM AP data.

**Methods**

**Modelling Ileak**

We added a leak equation to the Kernik and Paci IPS-CM and ToR-ORD adult cardiomyocyte models. Knowing that leak acts as a depolarizing current in
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For both models, decreases in \( R_m \) the perforating agent slowly decreased the access resistance to the cell. After contact was made with a cell and a seal of \( R_m \), a cell was used to estimate the electrical properties of the cell. We calculate access resistance \( R_a \) since it is equal to the parallel resistances of \( R_m \) and \( R_s \). The presence of \( R_m \) introduces uncertainty when \( R_m \) is used to approximate \( R_a \), making it difficult to accurately correct for leak current effects. For simplicity, we have omitted other elements of this patch clamp diagram (e.g. series resistance and capacitance).

\[
\text{Figure 1} \quad R_{\text{seal}} \text{ cannot be measured directly once access is gained. Once access is gained, we can only measure the combined resistance } R_m, \text{ which is equal to the parallel resistances of } R_m, \text{ and } R_s. \text{ The presence of } R_m, \text{ introduces uncertainty when } R_m, \text{ is used to approximate } R_a, \text{ making it difficult to accurately correct for leak current effects. For simplicity, we have omitted other elements of this patch clamp diagram (e.g. series resistance and capacitance).}
\]

\[ R_{\text{seal}} = \frac{1}{R_m} V = g_{\text{seal}} V. \tag{1} \]

where \( R_{\text{seal}} \) is the seal resistance and \( V \) denotes the membrane potential. The inverse of \( R_{\text{seal}} \) is the conductance, \( g_{\text{seal}} \). Note that more complicated equations for leak current (non-linear, and/or with a non-zero reversal potential) may be required in experiments where \( C_{\text{seal}} \) seal enhancer is used.\(^{24}\)

The effect of \( I_{\text{leak}} \) on the evolution of \( V \) was modelled as follows:

\[ \frac{dV}{dt} = -\frac{1}{C_m} (I_{\text{leak}} + I_{\text{ion}}), \tag{2} \]

where \( I_{\text{leak}} \) represents the sum of transmembrane currents and \( C_m \) is the membrane capacitance. \( C_m \) was set to 50 pF (the experimental average from the cells used in the present study) for the Kernik and Paci simulations, and for ToR-ORD, a value of 50 or 153 pF (the ToR-ORD baseline capacitance) was used unless specified otherwise.

Electrophysiological setup and data analysis

Perforated patch clamp experiments were conducted following a previously described protocol (see Supplementary Methods for more details).\(^{25}\)

After contact was made with a cell and a seal of >300 MΩ was formed, the perforating agent slowly decreased the access resistance to the cell (usually 10–15 min). This low \( R_m \) acceptance criterion was selected because we wanted to explore seal–leak effects above and below 1 GΩ. A series resistance \( R_s \) of 9–50 MΩ was maintained for all experiments. In this study, we used all cells from Clark et al.\(^{22}\) with membrane resistance \( R_m \) and \( R_s \) measurements acquired before and after current clamp recordings and that did not produce spontaneous alternans \((n = 37 \text{ out of } 40 \text{ cells})\). \( R_m \), \( C_m \), and \( R_s \) values were measured at 0 mV within 1 min prior to the acquisition of current clamp data.

All action potential (AP) features were calculated using a 10-s sample of current clamp data. The minimum potential \( V_{\text{cmd}} \) was measured twice during iPSC-CM experiments.\(^{20}\)

\( R_{\text{seal}} \) as an estimate of \( R_{\text{seal}} \)

We calculate \( R_{\text{seal}} \) using a small test pulse in voltage clamp mode.\(^{26}\)

\[ R_{\text{seal}} = \frac{\Delta V_{\text{cmd}}}{\Delta I_{\text{out}}}. \tag{3} \]

Here, \( \Delta V_{\text{cmd}} \) is the applied voltage step, and \( \Delta I_{\text{out}} \) is the difference in recorded current from before to during the step. Once access is gained to a cell, it can be difficult to estimate \( R_{\text{seal}} \), as the measured input resistance \( R_m \) depends on both \( R_m \) and \( R_{\text{seal}} \). The effect of patch clamp series resistance on \( R_m \) was excluded from Eq. (4).

\[ 1/R_m = 1/R_m + 1/R_{\text{seal}} \tag{4} \]

The smallest \( R_{\text{seal}} \) considered was 300 MΩ, whilst \( R_s \) values ranged from 9 to 50 MΩ. An increase of \( R_s \) from 9 to 50 MΩ (a worst-case scenario we never observed) for a cell with a 300 MΩ \( R_{\text{seal}} \) would change \( R_m \) by 13%.

So, whilst \( R_m \) can change in these experiments, it is unlikely to affect \( R_m \) by more than a few percent, and \( R_{\text{seal}} \) is likely the predominant parameter affecting changes of \( R_{\text{seal}} \).

Additional methods

Additional methods can be found in the Supplementary material.

Results

Leak affects human-induced pluripotent stem cell-derived cardiomyocytes action potential morphology even at seal resistances above 1 GΩ

To investigate the effects of leak current on AP morphology, we simulated the addition of \( I_{\text{leak}} \) in the Kernik\(^{12}\) and Paci\(^{24}\) iPSC-CM models (Figure 2). Simulated AP recordings show that \( I_{\text{leak}} \) substantially alters AP morphology, even when \( R_{\text{seal}} \geq 1 \text{ GΩ} \), a common threshold used in cardiac patch clamp experiments.\(^{20}\) For both models, decreases in \( R_{\text{seal}} \) depolarize the MP and cause a decrease in the \( dV/dt_{\text{max}} \) likely due to an incomplete recovery of sodium channels at these depolarized MPDs. Indeed, the Kernik model shows a transition to a small oscillation with very low upstroke velocity when \( R_{\text{seal}} < 3 \text{ GΩ} \) and then depolarized quiescence when \( R_{\text{seal}} < 2 \text{ GΩ} \). \( I_{\text{leak}} \) effects on the APD\(_{90} \) differ for the two models—decreases to \( R_{\text{seal}} \) cause AP prolongation in the Paci model and AP shortening in the Kernik model. There are also differences in the effect of \( R_{\text{seal}} \) on CL: in the Kernik model, decreases in \( R_{\text{seal}} \) lead to a gradual decrease in CL, whilst in the Paci model, decreasing \( R_{\text{seal}} \) initially has limited effect on CL but then causes shortening as \( R_{\text{seal}} \) decreases below 5 GΩ.

Leak effects on adult cardiomyocyte action potentials are moderated by different current densities and increased ionic currents

The ToR-ORD adult cardiomyocyte model is also susceptible to \( I_{\text{leak}} \) effects, but the extent depends on cell capacitance (Figure 3). Simulations with \( C_m \) set to the average iPSC-CM capacitance (50 pF) result in substantial AP morphological changes when \( R_{\text{seal}} \) is between 1 and 2 GΩ. However, when \( C_m \) is set to a value in the range of adult human ventricular cardiomyocytes (153 pF), \( I_{\text{leak}} \) has little effect on AP morphology when \( R_{\text{seal}} \geq 1 \text{ GΩ} \) (Figure 3B).

\( R_{\text{seal}} \) is not stable

Unlike voltage clamp recordings, the effects of \( I_{\text{leak}} \) on AP morphology (measured in current clamp mode) cannot be corrected in post-processing. Current clamp leak compensation is a potential solution to the issue\(^{22,23}\) but requires an accurate measure of \( R_{\text{seal}} \) throughout the experiment. \( R_{\text{seal}} \) cannot be accurately determined after access is gained because measures are contaminated by \( R_m \); such resistance measures are a composite of these two resistances that we nominally refer to as \( R_m \) (see Figure 1 and Methods). It is, therefore, tempting to measure the value before gaining access and assume it remains unchanged for the duration of an experiment. To investigate this, we considered \( R_{\text{seal}} \) measures taken two times during iPSC-CM experiments. \( R_m \) was measured with 5 mV steps from a holding potential of 0 mV (i.e. the leak reversal

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potential) before and after acquiring current clamp data. The data are skewed, with a mean of $R_{in} = 2.71 \Omega$ and median of $R_{in} = 0.82 \Omega$.

The relative change in $R_{in}$ from the first to the second time point was calculated and is plotted against the time elapsed between $R_{in}$ measurements in Figure 4B. The median change of $R_{in}$ is −15%. Because positive and negative changes cancel each other out in these statistics, we also inspected the absolute change, where we found a median of 20%. These data illustrate that $R_{seal}$ measurements often change over time. If we assume $R_{seal}$ is stable during experiments, this change in $R_{in}$ should be attributed to $R_{seal}$ and suggests that the average cell’s $R_{seal}$ decreases (and therefore $I_{leak}$ increases) over time.

**R**$_{in}$ is not a good approximation of $R_{seal}$ at any holding potential

A holding potential of −80 mV is a common choice for approximating $R_{seal}$ with $R_{in}$ measures. At this potential, sodium, calcium, and several potassium currents are expected to be largely inactive, but contributions from both $I_{K_1}$ and $I_f$ must still be considered. Whilst $I_{K_1}$ is perhaps close to its reversal potential (and therefore small), $I_f$ is not and can play a large role at this voltage.

We recently showed that $I_f$ is present in at least some of the iPSC-CMs used in this study. $I_f$ is also present in both the Kernik and Paci models, and we found the dynamics of the Kernik $I_f$ model to be quite similar to the in vitro data in this study (Figure 5A and B). Figure 5A shows an example cell’s response to an $I_f$-activating hyperpolarizing step before and after treatment with quinine, at a concentration expected to lead to 32% $I_f$ block (these data are taken from a section of a larger protocol—see Clark et al. Figure 6A). A change in total current of nearly 2 A/F is observed after hold at −120 mV for 1 s (Figure 5A). In Clark et al., nine cells were treated with quinine, and the average change during the $I_f$-activating segment was 1.34 A/F. We found that these nine cells could be sorted into three triplets based on the amount of quinine-induced $I_{leak}$ change during the $I_f$ segment: no/little sensitivity ($\Delta I_{leak}$ of 0–0.2 A/F), moderate sensitivity ($\Delta I_{leak}$ of 0.7–1.2 A/F), and large sensitivity ($\Delta I_{leak}$ of >1.9 A/F). Simulations using the Kernik model with 32% block of $I_f$ show a change of 1 A/F (i.e. moderate change) in $I_{leak}$ (Figure 5B).

To illustrate the effect of $I_f$ on leak calculations, we compared simulations from Kernik + leak models with $R_{seal} = 1 \Omega$ and with $g_f$ set to zero (i.e. not sensitive to quinine during hyperpolarizing step), the Kernik baseline value ($g_f = 0.0435 \text{ nS/pF}$, i.e. moderate sensitivity), or twice its baseline value ($g_f = 0.087 \text{ nS/pF}$, i.e. large sensitivity).
Figure 3  Effect of $R_{\text{seal}}$ on ToR-ORd adult cardiomyocyte APs at 50 and 153 pF. Simulations from the ToR-ORd + leak model paced at 1 Hz with $C_m$ set to 50 (A) and 153 pF (B), and $R_{\text{seal}}$ set to values from 1 to 10 GΩ. The dashed (red) trace shows a baseline (leak-free) simulation. Three AP morphology metrics for the 50 and 153 pF models are plotted against $R_{\text{seal}}$ (displayed on log-scaled x-axis): APD$_{90}$, action potential duration at 90% repolarization; $dV/dt_{\text{max}}$, maximum upstroke velocity; MP, minimum potential.

Figure 4  $R_{\text{in}}$ changes during iPSC-CM experiments. (A) Distribution of initial $R_{\text{in}}$ measurements from iPSC-CMs acquired with a +5 mV step from 0 mV. (B) The percentage change in $R_{\text{in}}$ plotted against the time elapsed between $R_{\text{in}}$ measurements. The interval between measurements ranged from 1 to 10 min. Time was recorded to the nearest minute, leading to the appearance of banding in the $\Delta$Time measure.
Figure 5 Ignoring the presence of \( I_l \) makes it impossible to accurately measure \( R_{\text{seal}} \) after gaining access. (A) Voltage clamp data acquired from an iPSC-CM before and after treatment with quinine, which is expected to block 32% of \( I_l \) at the concentration used. (B) Kernik model response at baseline and with 32% block of \( I_l \). (C) Kernik + leak voltage clamp simulations conducted with \( R_{\text{seal}} = 1 \, \Omega \), \( g_{f,K} \) reduced by 90%, and \( g_f \) set to 0 (solid line), 0.0435 (dotted line), or 0.087 nS/pF (dashed line). A voltage step from −80 to −75 mV was applied, as is commonly used to estimate \( R_{m} \). This \( R_{in} \) value is sometimes used to approximate \( R_{\text{seal}} \) when the holding potential is near −80 mV. The amplifier-measured (\( I_{\text{ion}} \)), total transmembrane (\( I_{\text{ion}} \)), and leak currents (\( I_{\text{leak}} \)) displayed. The \( R_{in} \) values calculated based on \( \Delta V_{\text{ion}} \) are 2.03, 1.50, and 1.16 \( \Omega \) for the 0, 0.0435, and 0.087 nS/pF simulations, respectively. (D) \( R_{in} \) values are plotted against holding potential for Kernik + leak models with \( R_{\text{seal}} = 1 \, \Omega \) and \( g_f \) equal to 0, 0.0435, or 0.087 nS/pF. The horizontal dotted line shows the true simulated \( R_{\text{seal}} \) value of 1 \( \Omega \).

(Figure 5C). We also reduced \( g_{f,K} \) in these models to 10% of the baseline value to highlight the effects of \( I_l \) on \( R_{in} \) measures independent of \( I_{\text{ion}} \). The calculated \( R_{in} \) values for these models at −80 mV are 2.03 \( \Omega \) for \( g_f = 0 \) nS/pF (little change), 1.50 \( \Omega \) for \( g_f = 0.0435 \) nS/pF (moderate change), and 1.16 \( \Omega \) for \( g_f = 0.087 \) nS/pF (large change) (Figure 5C). These simulations show that, at −80 mV, \( I_l \) contributes to \( I_{\text{leak}} \) and affects measures of \( I_{\text{seal}} \).

Using these same models, we then calculated \( R_{in} \) values at multiple holding potentials between −90 and +30 mV to determine whether we could find a potential where \( R_{in} \) is close to \( R_{\text{seal}} \), thereby minimizing the prediction error (Figure 5D). The model predicts that 20 mV (\( R_{in} = 0.96 \, \Omega \)) minimizes the error in our approximation of \( R_{seal} \). This does not mean that \( R_{in} \) measurements at 20 mV will always produce the best estimate of \( R_{\text{seal}} \). Instead, it indicates the size of \( I_{\text{leak}} \) does not change much when taking a 5 mV step from this potential. There is, however, a considerable amount of total current present, making this \( R_{\text{seal}} \) prediction sensitive to variations in the predominant ionic currents at this potential. Moreover, \( I_{\text{leak}} \) will be small and therefore more difficult to measure as 10 mV is close to the leak reversal potential (0 mV). It is also worth noting that the complex voltage-and time-dependent behaviour of transmembrane currents make \( R_{in} \) measures sensitive to both the duration and size of the voltage step (e.g., see supplement to Clerx et al.29). In summary, it is difficult to find a holding potential where \( R_{\text{seal}} \) can be measured without contamination from any transmembrane currents (i.e., where \( I_{\text{leak}} \approx I_{\text{ion}} \)).

Taken together, these findings provide evidence to the claim that \( R_{\text{seal}} \) cannot be reliably measured in iPSC-CMs once access is gained.

Next, we compared the effect of \( I_l \) on \( R_{in} \) and investigated the error in assuming \( R_{\text{seal}} \approx R_{in} \) at both a 0 mV (i.e., \( I_{\text{leak}} \) reversal) and −80 mV holding potential. At 0 mV, the Kernik + leak model is not sensitive to changes in \( g_f \), as \( I_l \) is largely non-conductive (Figure 6A). However, due
to an increased relative contribution of inward currents at 0 mV, the Kernik + leak model predicts a \( R_{\text{in}} \) with a large overestimation of \( R_{\text{in}} \) (Figure 6B). This error increases as the true value of \( R_{\text{in}} \) increases. Figure 6B also illustrates the sensitivity of the model to variations in \( g_f \) at \(-80\) mV, with \( R_{\text{in}} \) estimation errors decreasing as \( g_f \) increases; these errors also increase as \( R_{\text{in}} \) increases. The improved prediction accuracy of the 0.087 nS/pF model at \(-80\) mV is a coincidental side effect of doubling \( g_f \) with a different distribution of ion current densities or a larger baseline \( g_f \) value, the same doubling could just as easily worsen \( R_{\text{in}} \) predictions. For example, the \( g_f \) of an iPSC-CM with a large \( I_{\text{out}} \) current may slightly underestimate \( R_{\text{in}} \) at \(-80\) mV—doubling \( g_f \) in this case would result in a greater underestimation, increasing the error of the estimate.

### \( C_{\text{m}} \) and \( R_{\text{in}}(0\text{ mV}) \) correlate with minimum potential

The iPSC-CMs used in this study displayed a heterogeneous phenotype (Figure 7), producing both spontaneously firing (\( n = 25 \)) and non-firing (\( n = 12 \)) current clamp recordings. Figure 7A shows three cells with very different baseline current clamp recordings: non-firing and depolarized (green), spontaneously firing with a short AP (teal), and spontaneously firing with a long AP (red). Non-firing cells (MP = \(-42 \pm 8\) mV) and cells with spontaneously firing APs were depolarized (MP = \(-54 \pm 7\) mV)—the spontaneously firing cells also had a shorter AP duration (APD\(_{90} = 128 \pm 71\) ms) (Figure 7B) relative to adult cardiomyocytes and iPSC-CM models.13,14

We used linear regression analyses to determine if there is a correlation between \( g_{\text{in}}/C_{\text{m}} \) and AP biomarkers. Here, we use \( g_{\text{in}} \) (instead of \( R_{\text{in}} \)), as it reduces the spread of this variable and positively correlates with \( I_{\text{leak}} \) providing a more interpretable comparison with AP morphology. The values of each cell’s \( g_{\text{in}} \) and \( C_{\text{m}} \) are shown in Figure 7C. \( I_{\text{leak}} \)’s effect on AP morphology is expected to scale directly with \( g_{\text{in}} \) and inversely with \( C_{\text{m}} \). This is because \( g_{\text{in}} \), even if a poor estimate, is expected to correlate with \( g_{\text{leak}} \) (Figure 6B) A given \( g_{\text{leak}} \) will cause a smaller contribution in larger cells (i.e. cells with larger \( C_{\text{m}} \)), because the ionic currents are expected to scale with the size of the cell. For this reason, four AP biomarkers (MP, APD\(_{90}\), CL, and \( dV/dt_{\text{max}} \)) were compared with \( g_{\text{in}}/C_{\text{m}} \) (Figure 8). The MP of spontaneously firing (\( R = 0.44, P < 0.05 \)) and non-firing (\( R = 0.76, P < 0.05 \)) cells are positively correlated with \( g_{\text{in}}/C_{\text{m}} \) (Figure 8A). This finding is in agreement with our in silico studies showing that increasing \( g_{\text{leak}} \), thereby increasing \( g_{\text{in}} \), will depolarize the cell (Figure 7). The other three biomarkers failed at least one of the assumptions required when conducting a linear regression analysis (see Supplementary Methods). There are no obvious trends when comparing \( g_{\text{in}}/C_{\text{m}} \) with CL or \( dV/dt_{\text{max}} \). The APD\(_{90}\) plot, however, indicates there may be some AP shortening as \( g_{\text{in}}/C_{\text{m}} \) increases. Due to under-sampling and a lack of linearity, we cannot make any claims of significance between these two measures. Leak simulations with the models, though correlated, did not predict a linear relationship between \( g_{\text{leak}} \) and these biomarkers (Figure 2C and D). However, the MP vs. \( g_{\text{in}}/C_{\text{m}} \) relationship passes all tests of linear regression assumptions and trends in the same direction as the Kernik and Paci simulations in Figure 2.

### Fitting background currents in human-induced pluripotent stem cell-derived cardiomyocyte models can absorb and imitate \( I_{\text{leak}} \)

We used optimization to study the potential of linear background currents (e.g. sodium and calcium) to imitate leak effects (see Supplementary Methods). We fit the baseline Kernik model to a Kernik + leak model with \( R_{\text{in}} = 5 \text{ G}\Omega \) (Figure 9), allowing only the background sodium (\( g_{\text{Na}} \)) and background calcium (\( g_{\text{Ca}} \)) conductances to vary. These currents were selected because they were incorporated into the Kernik model without independent iPSC-CM experimentation or validation. The best-fit model had an increased \( g_{\text{Na}} \) (\( \times 7.0 \)), whilst \( g_{\text{Ca}} \) (\( \times 1.0 \)) did not change much relative to the baseline model (Figure 9A). Whilst not...
In this study, differences in the experiments, but not when cells were clustered.11,12

CL, even with that leak current substantially depolarizes the MP and shortens the

similar to adult human cells (with model

Leak affects action potential morphology

Leak current is a common and unavoidable experimental artefact that affects patch clamp recordings. In this study, using both model predictions and experimental data, we show that leak current: (i) affects iPS-CM AP morphology, (ii) can vary during experiments, (iii) cannot be accurately estimated after access is gained to an iPS-CM, and (iv) may be absorbed by linear equations for background currents when iPS-CM models are fit to experimental AP data. During iPS-CM current clamp studies, leak consideration often starts with a pre-rupture seal measurement (with a 1 GΩ threshold) and is ignored if the seal appears to remain stable throughout the study. Here, we argue leak effects should be quantitatively scrutinized during the acquisition, analysis, and fitting of experimental data. Furthermore, we believe cell-to-cell variation in seal resistance contributes to observed iPS-CM AP heterogeneity—often attributed nearly entirely to variations in ionic current densities.

A perfect match, the best-fit trace reproduced qualitative features of the baseline + leak trace, showing a depolarized MP and a smaller amplitude (Figure 9B). This indicates that increased Ileak can affect the AP in a fashion similar to Ileak such that mathematical iPS-CM models may absorb Ileak effects by erroneously increasing background currents.

Discussion

Leak current is a common and unavoidable experimental artefact that affects patch clamp recordings. In this study, using both model predictions and experimental data, we show that leak current: (i) affects iPS-CM AP morphology, (ii) can vary during experiments, (iii) cannot be accurately estimated after access is gained to an iPS-CM, and (iv) may be absorbed by linear equations for background currents when iPS-CM models are fit to experimental AP data. During iPS-CM current clamp studies, leak consideration often starts with a pre-rupture seal measurement (with a 1 GΩ threshold) and is ignored if the seal appears to remain stable throughout the study. Here, we argue leak effects should be quantitatively scrutinized during the acquisition, analysis, and fitting of experimental data. Furthermore, we believe cell-to-cell variation in seal resistance contributes to observed iPS-CM AP heterogeneity—often attributed nearly entirely to variations in ionic current densities.

Leak affects action potential morphology

Simulations in chick embryonic cardiomyocytes, which are smaller than adult human cells (with model Cm = 25.5 pF), have previously shown that leak current substantially depolarizes the MP and shortens the CL, even with Rseal values of 5 GΩ.29 More recently, it was shown that in vitro iPS-CMs were significantly depolarized during single-cell experiments, but not when cells were clustered.11,12 These results indicate that isolated iPS-CMs are likely affected by leak current. Our in vitro and in silico findings support this conclusion and strengthen the argument that iPS-CM AP morphology is strongly affected by leak current.

Our in silico work indicates that Ileak has a smaller effect on recordings of adult cardiomyocyte AP morphology when compared with iPS-CMs (Figure 3B). This effect is strongly modulated by Cm, indicating the larger size of adult cardiomyocytes has a moderating effect on Ileak-induced AP changes. When the Ileak artefact in this adult model is normalized by the average iPS-CM capacitance (50 pF, Figure 3A), Ileak substantially alters the AP shape at Rseal values above 1 GΩ. But the effects are much less than in the iPS-CM model (Figure 2)—this indicates the ionic current expression profile of adult cardiomyocytes (e.g. greater Ina and lower Ileak density) can vary from one cell to the next, and modulates the effects of Ileak on adult AP recordings. Thus, differentiation strategies that aim to mature the iPS-CM phenotype (both in size and ionic current expression) will likely produce cells that are affected less by Ileak artefact.

Human-induced pluripotent stem cell-derived cardiomyocytes have long been defined by their immature and heterogeneous electrophysiological phenotype.10,30 Such features are due, at least in part, to the types of ion channels expressed and cell-to-cell variations in ionic current conductances.10,30 In this study, differences in the Ileak responses to nine quinine-treated cells are an example of how iPS-CM ionic currents can vary from one cell to the next. Heterogeneity in AP morphology and ionic current expression is also seen in primary adult cardiomyocytes.31–33

In this study, we show that Ileak also contributes to this immature and heterogeneous AP phenotype during single-cell patch clamp experiments. The relative importance of Ileak’s influence on AP shape varies amongst cells and depends on several factors, including Rseal, Cm, and the ionic current expression profile. Simulations indicate that the AP shape can be substantially altered (relative to non-patched cells), even when Rseal is equal to 10 GΩ, an unrealistically high acceptance criterion for iPS-CM patch clamp studies. These factors, along with the potential for Rseal to change during an experiment, can confound drug and genetic mutation studies. For example, the irregular and...
depolarized phenotype (caused at least in part by \( I_{\text{leak}} \)) of iPSC-CMs in our recent cardiotoxicity study\(^{25} \) made it impossible to measure consistent cell-specific changes in spontaneous AP morphology from pre- to post-drug application.

The AP-altering effects of \( I_{\text{leak}} \) can be effectively eliminated by patching cells whilst in engineered heart tissue or monolayer. The electrical coupling of cells in these conditions results in an enormous effective capacitance, rendering \( I_{\text{leak}} \) an infinitesimal contributor to total current. Whilst this eliminates the \( I_{\text{leak}} \) artefact, it also comes at a cost—this approach does not allow for the direct measure of APs in individual cells, limiting the ability to study iPSC-CM heterogeneity. In addition, it is not possible to acquire voltage clamp data from cells in these conditions—as such, one could not acquire both AP and descriptive data about individual currents, as we recently have done in isolated cells.\(^{25} \)

**Predicting \( R_{\text{seal}} \) during experiments**

\( R_{\text{seal}} \) can be well approximated prior to gaining access to a cell, but after perforation (or rupture), the presence of membrane currents makes it impossible to obtain an accurate measurement (Figure 5). Our *in silico* work shows that, even when currents such as \( I_{f} \) and \( I_{K1} \) are reduced to <10% of their baseline values, \( R_{m} \) (measured at −80 mV) is still a poor approximation of \( R_{\text{seal}} \) (Figure 6, solid black line).

To address these difficulties, we believe it may be feasible to use the pre-rupture \( R_{\text{seal}} \) and post-rupture \( R_{m} \) measures to calculate estimates of \( R_{\text{seal}} \) during an experiment. This approach would require an accurate measure of \( R_{m} \) just after access is gained. Using \( R_{\text{seal}} \) and the initial \( R_{m} \), it is possible to calculate \( R_{m} \) (Figure 1). An estimate of \( R_{\text{seal}} \) could then be made at any time during the experiment, assuming the calculated \( R_{m} \) stays constant, by re-measuring \( R_{m} \) and using Eq. (4). This approach relies on two major assumptions: (i) the perforation/rupture step does not affect the seal, and (ii) a protocol or procedure exists that can be used prior to each measurement of \( R_{m} \) to ensure that the contribution of \( R_{m} \) is consistent. We cannot say for certain that these assumptions will always be valid. However, we believe that recording frequent \( R_{m} \) measurements, estimating \( R_{\text{seal}} \), and scrutinizing changes are important steps for the correct interpretation of iPSC-CM current clamp data.

**Correcting for \( R_{\text{seal}} \) during experiments**

We believe these \( R_{\text{seal}} \) estimates should be used in a dynamic clamp leak compensation setup to address the limitations caused by a depolarized

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**Figure 8** Relationship between \( g_{\text{in}}/C_{m} \) and AP biomarkers. (A) \( g_{\text{in}}/C_{m} \) plotted against MP. Spontaneously firing cells are denoted as teal circles and non-firing cells as red squares. Linear regression fits to data from spontaneous (teal dashed, \( R = 0.47, P < 0.05 \)) and non-firing (red dotted, \( R = 0.76, P < 0.05 \)) cells are overlaid on the plot. No statistically significant relationship was found between \( g_{\text{in}}/C_{m} \) and APD\(_{90} \), CL (C), or \( dV/dt_{\text{max}} \) (D). APD\(_{90} \), action potential duration at 90% repolarization; CL, cycle length; \( dV/dt_{\text{max}} \), maximum upstroke velocity; MP, minimum potential.
Models of background currents can incorporate leak artefacts

The Kernik and Paci iPSC-CM models took ion-specific background currents from the ten Tusscher et al. model. These currents can trace their roots to the seminal work of Luo et al. where they were included to help maintain physiologically realistic intra-cellular concentrations.

Direct measurements of \( I_{bNa} \) and \( I_{bCa} \) in iPSC-CMs have not been reported. The Kernik and Paci iPSC-CM models both adopted the ventricular formulation for \( I_{bCa} \) and \( I_{bNa} \) and then set the conductances of these currents by comparing model predictions of the AP with in vitro measurements in iPSC-CMs. We posit that \( I_{bNa} \) is overestimated and compensates for the explicit consideration of leak current artefacts, a source of discrepancy between these models and reality. We expect consideration of leak when constructing iPSC-CM models to reduce background sodium current and result in a more realistic model of intact iPSC-CMs.

Modelling experimental artefacts

Whilst the effects of experimental artefacts in single-cell studies are well-established, consideration of them whilst building ion channel and AP models has been limited. In silico studies investigating series resistance effects on voltage clamp recordings have been done in fast-activating currents, such as \( I_{Na} \) and \( I_{hp} \), but to our knowledge, artefact equations have not been included in the calibration process for widely used models of these currents—although the \( I_{hp} \) model by Ebihara et al. was incorporated directly into the widely copied \( I_{Na} \) model by Luo et al. Recently, Lei et al. demonstrated that coupling experimental artefact equations with an \( I_{hp} \) mechanistic model improved predictions. These studies show that including experimental artefact equations in model fitting can improve the descriptive ability of the resulting electrophysiological models. As such, we believe experimental artefacts should be explicitly considered at the modelling phase and not ignored simply because a pre-determined minimum threshold is reached (e.g. 1 GΩ). Based on our findings, we believe cardiomyocyte models and especially iPSC-CM models should explicitly include leak currents when fitting to experimental current clamp data.
**Recommendations**

Our results provide important insights and recommendations for experimentalists and modellers alike:

1. **Experimental:** $R_{\text{leak}}$ should be recorded before gaining access to a cell and $R_m$ measured frequently during an experiment. It is important to measure $R_m$ from a voltage that provides a consistent measure of $R_{\text{leak}}$ such that any changes in $R_m$ can be attributed to changes in $R_{\text{leak}}$.

2. **Experimental:** Dynamic injection of a leak compensation current can help a cell recover its native AP, including the MP. Because $R_{\text{leak}}$ is difficult to measure during experiments and to avoid over-compensation, we advise under-compensation (e.g. 70%). Additionally, $R_{\text{leak}}$ and $R_m$ measurements should be reported.

3. **Modelling:** Explicit inclusion of $I_{\text{leak}}$ will improve the descriptive ability of iPSC-CM models. Whilst this may not always improve fits to AP data, it will take into account an important current affecting iPSC-CM recordings.

**Limitations and future directions**

This study has several limitations that should be considered during future investigations that may be affected by $I_{\text{leak}}$. First and foremost, when gathering these data for a previous study, we did not follow our new recommendation of recording the exact value of $R_{\text{leak}}$ before gaining access and then measuring $R_m$ just after perforation. Going forward, we hope to use these two values to predict $R_{\text{leak}}$ at multiple time points during an experiment, as outlined in Section 3.2. Second, we only conducted these experiments in one cell line. Whilst our results appear similar to data from other labs, it would be useful to conduct this study on multiple cell lines in the same lab. Third, we did not attempt dynamic injection of a leak compensation current—in future work, we would like to investigate this as an approach to reducing cell-to-cell heterogeneity. Finally, the iPSC-CM models have innumerable differences from the cells used in this study, which is evident when comparing AP morphologies of in vitro cells (Figure 7A) to in silico models (Figure 2). However, the agreement that we did see between simulations and our in vitro data demonstrates the potential of improving the descriptive ability of iPSC-CM models by including a leak current.

**Conclusion**

In this study, we demonstrate that leak current affects iPSC-CM AP morphology, even at seal resistances above 1 GΩ, and contributes to the heterogeneity that characterizes these cells. Using both in vitro and in silico data, we showed the challenges of estimating $R_{\text{leak}}$ after gaining access to a cell and that $R_{\text{leak}}$ is subject to change during the course of an experiment. We also posit that background sodium current in both steps of the way.

**Supplementary material**

Supplementary material is available at [Europeonline](https://www.europeonline.com).

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**Data availability**

All data, code, and models can be accessed from GitHub ([https://github.com/Christini-Lab/iPSC-leak-artifact](https://github.com/Christini-Lab/iPSC-leak-artifact)).

**Translational perspective**

Human iPSC-CMs have emerged as a promising translational tool to study human cardiac physiology outside of the clinic. They have been particularly useful to investigate cell-level pro-arrhythmic substrates, including genetic mutations and ion channel-blocking drugs, and play a critical role as a model for validating drug effects on human whole-cell electrophysiology in the Comprehensive in vitro Proarrhythmia Assay (CiPA) initiative. However, the depth of insights from iPSC-CM data is often limited by inter- and intra-lab heterogeneity, caused at least in part, by patch clamp experimental artefact. In this manuscript, we show how the seal–leak current is an often-overlooked artefact that confounds studies with iPSC-CMs. Ultimately, the findings and recommendations within this manuscript will improve the use of iPSC-CMs as an in vitro model to study cardiac electrophysiological diseases and patient-specific treatment strategies.

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