Extracellular Charge Adsorption Influences Intracellular Electrochemical Homeostasis in Amphibian Skeletal Muscle

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ABSTRACT The membrane potential measured by intracellular electrodes, $E_m$, is the sum of the transmembrane potential difference ($E_1$) between inner and outer cell membrane surfaces and a smaller potential difference ($E_2$) between a volume containing fixed charges on or near the outer membrane surface and the bulk extracellular space. This study investigates the influence of $E_2$ upon transmembrane ion fluxes, and hence cellular electrochemical homeostasis, using an integrative approach that combines computational and experimental methods. First, analytic equations were developed to calculate the influence of charges constrained within a three-dimensional glycocalyx matrix enveloping the cell membrane outer surface upon local electrical potentials and ion concentrations. Electron microscopy confirmed predictions of these equations that extracellular charge adsorption influences glycocalyx volume. Second, the novel analytic glycocalyx formulation was incorporated into the charge-difference cellular model of Fraser and Huang to simulate the influence of extracellular fixed charges upon intracellular ion homeostasis. Experimental measurements of $E_m$ supported the resulting predictions that an increased magnitude of extracellular fixed charge increases net transmembrane ionic leak currents, resulting in either a compensatory increase in Na$^+$/K$^+$-ATPase activity, or, in cells with reduced Na$^+$/K$^+$-ATPase activity, a partial dissipation of transmembrane ionic gradients and depolarization of $E_m$.

INTRODUCTION

The electrical potential difference between the intracellular and bulk extracellular spaces ($E_m$), as classically measured by intracellular microelectrodes, is thought to consist of at least two distinct components, as shown in Fig. 1, such that $E_m = E_1 + E_2$. The larger $E_1$ term reflects the potential difference between the inner and outer membrane surfaces. The smaller $E_2$ term is the potential difference between regions containing fixed negative charges on or near the outer membrane surface, and the bulk extracellular space. A further potential difference between the inner membrane surface and the bulk intracellular space is also likely, but would be constant in the absence of large alterations in intracellular fluid composition (1). Previous investigations of the physiological effects of alterations in $E_2$ by Ca$^{2+}$ adsorption onto these fixed negative charges (2,3) demonstrated alterations in ion channel function at a constant $E_m$ that suggested they were gated by $E_1$ rather than by $E_m$ itself (1,3,4). However, these previous studies did not explore the possible consequences of alterations in $E_2$ upon electrophysiological homeostasis in intact whole cells, instead investigating isolated ionic currents studied under voltage clamp.

This study thus extends and complements the earlier works through an investigation of the influence of $E_2$ upon transmembrane ionic currents and intracellular ion homeostasis. Computational methods were used to develop testable theoretical predictions derived from the charge-difference (CD) approach (5) previously used to investigate the determinants of intracellular ion concentrations, membrane potential, and cell volume in complex multicompartment systems involving both passive and active ion fluxes (5–8). The theoretical predictions were tested against focused experimentation in frog skeletal muscle.

First, the existence of a discrete, negatively charged, glycoproteaceous glycocalyx known to possess Ca$^{2+}$-adsorptive properties surrounding skeletal muscle prompted the development of a mathematical description of the influence of fixed charges distributed within a three-dimensional volume. Electron microscopy showed the glycocalyx to be 35–70 nm thick in this study, one to two orders of magnitude greater than the Debye length calculated for the Ringer solutions employed. This theoretical analytic approach was developed from Gibbs-Donnan (GD)-type equilibrium constraints that simulated the effects of alterations in the magnitude of fixed glyocalyx charge within a three-dimensional volume. This required consideration of whether osmotic alterations in glyocalyx volume would influence the relationship between glyocalyx charge adsorption, $E_2$, and local ion concentrations. Electron micrographic measurements demonstrated that the thickness of the glycocalyx was influenced by extracellular [Ca$^{2+}$], in agreement with predictions from a GD analysis that assumed constant osmolyte concentration within the glycocalyx. This suggested a nonlinear relationship between glyocalyx [Ca$^{2+}$] adsorption and steady-state glyocalyx charge density, in contrast to the linear relationship predicted by the classical Gouy-Chapman (GC) equation (9). The GC equation requires the assumption of a...
This study thus demonstrates an influence of glycocalyceal charge upon transmembrane ionic leak currents. Consistent with assumptions made in earlier studies, this had little influence upon $E_m$ under normal physiological conditions due to a compensatory increase in $\text{Na}^+/\text{K}^+$-ATPase activity. However, under conditions of compromised $\text{Na}^+/\text{K}^+$-ATPase activity produced by reduced extracellular $[\text{K}^+]$, glycocalyceal charge was shown to influence intracellular electrolyte homeostasis and $E_m$. These findings lead to a prediction that clinical conditions associated with reduced $\text{Na}^+/\text{K}^+$-ATPase activity, such as ischemia, hypokalemia, and cardiac failure (12), might also be exacerbated by reduced extracellular $[\text{Ca}^{2+}]$.

**MATERIALS AND METHODS**

Analytic equations based on GD theory were derived to assess the influence of extracellular fixed charges upon $E_2$. This approach was compared to the classical GC equations. Simulation of the physiological effects of $E_2$ upon intracellular ion homeostasis employed the CD approach, introduced and fully described on earlier occasions (5,7,8), and modified to include the GD equations. Finally, the related experimental studies involved conventional microelectrode measurements of membrane potential, and electron-microscope studies of glycocalyceal thicknesses. The symbols used in the text are summarized in Table 1.

**Experimental methods**

**Membrane potential recordings**

Resting membrane potentials, $E_m$, were recorded in sartorius muscles dissected from cold-adapted winter frogs, *Rana temporaria*, killed by concuss-
sion followed by pithing (Schedule 1, Animal Procedures Act, Home Office, United Kingdom). The frogs had been maintained together to minimize any variations that might reflect differing environmental influences. The muscles were stretched to ~1.5 times their in situ length as previously described by Koutsis et al. (13) and placed in a bath initially filled with standard isotonic Cl−-Ringer solution titrated to pH 7.2 (in mM: 115 NaCl, 2.5 KCl, 1.8 CaCl2, and 3.0 HEPES).

Experiments were conducted at 5–10°C. Bath temperature was controlled by circulating cooled water through a glass coil placed in the chamber using a Minipuls 3 peristaltic pump (Gilson, France) and monitored using a digital thermometer incorporating a remote thermometer close to the muscle, previously linearized and calibrated against a platinum film resistor. E<sub>m</sub> values from surface fibers were measured against bath voltage using standard glass capillary microelectrodes (3 M KCl; resistance 5–20 MΩ; tip potential <5 mV (14)) mounted through an Ag/AgCl reference junction to a high impedance voltage follower and controlled using a micromanipulator. Micro-electrode resistances were further tested after impaling each muscle fiber, and pedance voltage follower and controlled using a micromanipulator.

Experimental recordings of the influence of the experimental solutions upon E<sub>m</sub> were pooled from a time period (150–300 min) that under each experimental condition permitted detection of small changes in E<sub>m</sub> without visible deterioration or twitching of the muscle fibers under light microscopy.

Electron microscopy

Cutaneous pectoris muscles were obtained for electron microscopy from the same frogs used in the microelectrode experiments and exposed to the experimental condition. The glycocalyx will be assumed to influence the equilibrium state through its charge density. Mathematical analysis of the relationship between glycocalyceal fixed charge density, Σ<sub>E</sub>, glycocalyceal potential, E<sub>2</sub>, and local ion concentrations may be derived from two basic steady-state constraints: 1) electroneutrality, and 2) absence of net ion currents. First considering the situation in which Σ<sub>E</sub> is known, the electroneutrality constraint may be stated as

\[ Σ_E + F(\frac{[A]_g}{[B]_g}) = 0, \]  \hspace{2cm} (1)

where Σ<sub>E</sub> is the fixed charge/unit volume (C l<sup>-1</sup>); F is Faraday’s constant; A represents a monovalent cation; B represents a monovalent anion; and the subscript g denotes the glycocalyx. At equilibrium, ionic currents are zero and thus ions are passively distributed between the bulk extracellular solution and the glyocalyx according to their Nernst potentials:

\[ \frac{RT}{F} \ln \frac{a_a[A]_g}{a_B[B]_g} = E_2, \]  \hspace{2cm} (2)

where \( a_a \) means denote ionic activity in the extracellular fluid (ECF) and glyocalyx compartments, denoted by the subscripts e and g, respectively. This allows for the possibility that the glycocalyx may alter local ionic activities. This relationship may be simplified by defining [C] as the total salt concentration such that [C]<sub>E</sub> = [A]<sub>e</sub> = [B]<sub>e</sub>. Thus,

\[ [B]_g = \left( \frac{a_a}{a_B} \right) e^{(\frac{F}{RT})[C]_e}, \quad [A]_g = \left( \frac{a_a}{a_B} \right) e^{(\frac{F}{RT})[C]_g}. \]  \hspace{2cm} (3)

Combining these two constraints (Eqs. 1 and 3) demonstrates the relationship between fixed charge density and glyocalyceal potential.

\[ Σ_E = -F \left( \frac{a_a}{a_B} \right) \left( e^{(\frac{F}{RT})[C]_g} - e^{(\frac{F}{RT})[C]_e} \right). \]  \hspace{2cm} (4)

Rearranging in terms of E<sub>2</sub>,

\[ \frac{Σ_E}{2F} = \left( \frac{a_a}{a_B} \right) \frac{E_2}{RT} \arcsinh \left( \frac{a_B Σ_E}{2a_a F[C]_e} \right). \]  \hspace{2cm} (5)

This equation allows calculation of E<sub>2</sub> if the glycocalyx fixed charge density is known. However, if the glyocalyx volume is not fixed, but rather determined by osmotic water movements, then there must also be an osmotic equilibrium at steady state. Under such conditions, the relationship between total glyocalyx charge and the glyocalyx charge density may be nonlinear, as greater charge magnitudes result in an increase in the volume over which they distribute. This would have implications for the relationship between Ca<sup>2+</sup> adsorption and glyocalyx charge density, and thereby for E<sub>2</sub>. This analysis, therefore, will next develop an analytic relationship between glyocalyx charge and E<sub>2</sub> under conditions where glyocalyx volume is osmotically determined. The predictions of this theory will be compared with those of the nonsomatic relationship (Eq. 5) by electron-microscopic measurement of glyocalyx volumes under conditions of varying [Ca<sup>2+</sup>]<sub>e</sub>, in Results.

Rearranging Eq. 2 gives

\[ a_B[B]_g = \left( \frac{a_a}{a_B} \right) e^{(\frac{F}{RT})[B]_g}. \]  \hspace{2cm} (6)

The glycocalyx will be assumed to influence the equilibrium state through its charge and through its osmotic activity. Such influence is captured by defining Σ<sub>E</sub> as:

\[ Σ_E = Fz_{X(g)}[X]_g, \]  \hspace{2cm} (7)

where [X]<sub>E</sub> denotes the osmolar concentration of fixed ions within the glyocalyx and z_{X(g)} its mean charge valency per osmole. Note that this
permits $z_{X(g)}$ to be very large if the glycocalyx is significantly charged but has little osmotic activity. Thus, the electroneutrality constraint (Eq. 1) will be restated as

$$[A]_g - [B]_g + z_{X(g)}[X]_g = 0. \quad (8)$$

Finally, the osmotic constraint is given by

$$\phi_0[A]_g + \phi_0[B]_g + [X]_g = \eta \Pi_e, \quad (9)$$

where $\phi_0$ is the osmotic coefficient of the salt solution within the glycocalyx, $\Pi_e$ is the total osmolarity of the extracellular solution, and $[X]_g$ is expressed in osmoles $^{-1}$. Osmotic and chemical potentials alone may be shown to produce intrinsic instability in polyelectrolyte gels such as the glycocalyx, unless stabilized by a third force that may be considered to be a gel pressure (15). Thus, a dimensionless correction factor, $\eta$, is included in this osmotic constraint to reflect the possibility that total osmotic activity may be greater within the glycocalyx than in the ECF due to a small pressure within the glycocalyx. It will be assumed that $\eta$ is constant with changing glycocalyx volume within the range of values considered in this work.

Substituting the electroneutrality expression for $[A]_g$ (Eq. 8) into the Donnan constraint (Eq. 6) gives

$$a_e^2[B]_g - z_{X(g)}[X]_g a_e^2[B]_g - a_e^2[C]_e = 0. \quad (10)$$

Substituting Eq. 8 into the osmotic constraint (Eq. 9) gives

$$[B]_g = \frac{1 - \phi_0 z_{X(g)}}{-2 \phi_0}. \quad (11)$$

Combining these equations, grouping terms into quadratic form, and solving for $[X]_g$, yields

$$[X]_g = \frac{\eta \Pi_e \pm \phi_0}{1 - \phi_0^2 z_{X(g)}} \left( \frac{\eta \Pi_e z_{X(g)}^2}{2} + 4(1 - \phi_0^2 z_{X(g)}) a_e^2[C]_e \right). \quad (12)$$

This permits the expression of $E_2$ in terms of the mean charge valency per osmole of molecules constrained within the glycocalyx ($z_{X(g)}$), rather than in terms of the steady-state charge density within the glycocalyx ($\Sigma_g$). Thus, by combining Eqs. 5, 7, and 12, $E_2$ is shown to vary nonlinearly with $z_{X(g)}$:

$$E_2 = \frac{RT}{F} \arcsinh \left( \frac{z_{X(g)} a_e}{\eta \Pi_e \pm \phi_0} \left( \frac{\eta \Pi_e z_{X(g)}^2}{2} + 4(1 - \phi_0^2 z_{X(g)}) a_e^2[C]_e \right) \right). \quad (13)$$

Alternatively, if fixed charges in the glycocalyx itself exert no osmotic activity, then combining Eqs. 1, 6, and 9 with $[X]_g = 0$ gives

$$\Sigma_g = \left( \frac{\eta \Pi_e}{\phi_0} \right) \left( \frac{2 a_e^2[C]_e}{a_e^2} \right). \quad (14)$$

Thus, in the particular case that the glycocalyx itself exerts no osmotic influence but its volume is nevertheless osmotically determined,

$$E_2 = \frac{RT}{F} \arcsinh \left( \frac{a_e \sqrt{\frac{\eta \Pi_e}{\phi_0} - \frac{a_e^2[C]_e}{a_e^2}}}{2 a_e[C]_e} \right). \quad (15)$$

The relative volume of the glycocalyx at steady state is given by

$$V_{i(t \to \infty)} = \frac{[X]_{g(t \to 0)}}{[X]_{g(t \to \infty)}}. \quad (16)$$

or, equivalently,

$$V_{i(t \to \infty)} = \frac{\Sigma_g(t \to 0)}{\Sigma_g(t \to \infty)}, \quad (17)$$

where $V_i$ is the relative volume of the glycocalyx and times $t = 0$ and $t = \infty$ denote initial conditions and eventual steady-state conditions, respectively.

By inspection, it is clear that both Eqs. 12 and 14 have values of zero when $\Pi_e = 2 \phi_0 [C]_e$, $\eta = 1$, and $a_e = a_e$. The condition $\Pi_e = 2 \phi_0 [C]_e$ is likely to be true when the extracellular solution is a 1:1 electrolyte. Yet, as can be seen from Eq. 16 or Eq. 17, if steady-state values of $[X]_g$ or $\Sigma_g$ are zero, the steady-state glycocalyx volume tends toward “infinity” as the glycocalyx is then essentially indistinguishable from the bulk ECF. However, these equations additionally demonstrate that a stable, osmotically determined glycocalyx volume would be possible 1), if total osmolarity in the glycocalyx is permitted to be higher than in the ECF, thus $\eta > 1; 2)$, if the activities of mobile ions are greater in the glycocalyx than in the ECF, thus $a_e/a_e < 1; 3)$, if osmotic coefficients of mobile ions are lower in the glycocalyx than in the ECF, thus $\phi_0 < 1$ for ions within the glycocalyx such that $\Pi_e > 2 \phi_0 [C]_e$; or 4), if the ECF contains ions that cannot pass into the glycocalyx such that $\Pi_e > 2 \phi_0 [C]_e$. One or more of these conditions must be true for there to exist a stable, nonzero value of $[X]_g$ or $\Sigma_g$ if glycocalyx volume is osmotically determined. Certain other biological gels, including the sclera, have been shown to behave as if they exert a mechanical pressure that opposes swelling (15), and this (condition 1 above) seems the most likely explanation of the above four possibilities. Thus, unless otherwise stated, it is assumed that $a_e/a_e = 1$ and $\phi_0 = 1$. The influence of the value of $\eta$ was investigated, as shown in Results. Nevertheless, it is clear that any combination of the four possible conditions for stability presented here lead to the conclusion that if the glycocalyx volume is osmotically determined, then increasing the magnitude of $z_{X(g)}$ would tend to swell the glycocalyx. This hypothesis was tested experimentally:

Note that it is not necessary to consider the shape of the potential decay between the glycocalyx and the bulk ECF (Fig. 1). It can be shown that the electrochemical steady state of the glycocalyx is uniquely defined by the state of the ECF irrespective of whether intervening compartments are present. Thus, assuming activity and osmotic coefficients of 1, the steady-state value of $[X]_g$ (Eq. 12) is influenced by $[C]_e$ and $\Pi_e$ but not by any other property of the ECF. Yet by the osmotic and zero-net-flux constraints, $\Pi_e$ and $[C]_e$ must be the same in every linked compartment between which ions are passively distributed. Thus, the presence of intervening compartments does not alter the steady state of the glycocalyx with respect to the ECF, as $\Pi_e$ and $[C]_e$ must, by the same argument, be the same in each such compartment. Similarly, because activity ratios and osmotic coefficients can be expressed as ratios for ions in the glycocalyx relative to those in the ECF, this analysis holds true for situations where activity coefficients and osmotic coefficients vary between the glycocalyx edge and the ECF.

**Charge-difference modeling of the influence of $E_2$ upon intracellular ionic homeostasis**

The influence of glycocalyx ion concentrations and $E_2$ upon transmembrane ion fluxes and intracellular ionic homeostasis were explored using the CD model of Fraser and Huang (5,7,8). This employed an assumption that...
transmembrane ion movements were orders of magnitude slower than the passive redistribution of ions between the bulk extracellular space and the charged glycocalyx. This permitted analytic calculation of glycocalyx ion concentrations and \( E_2 \) from \( \Sigma_g \) and extracellular conditions using the GD method derived above.

CD modeling of the cell then introduced the assumption that transmembrane fluxes were determined by the electrical and respective concentration gradients between the bulk intracellular space and the glycocalyx. This contrasts with previous studies using this model in which transmembrane fluxes were determined by the concentration gradient between the bulk intracellular and bulk extracellular spaces. Extracellular ion concentrations were identical to those in the experimental solutions used. In other respects, the model was as described in Fraser and Huang (5), with the osmotic water movement terms later introduced by those authors (8), and therefore the membrane permeabilities and Na\(^+/K\(^{-}\)-ATPase density were calibrated to frog skeletal muscle.

RESULTS

Previous studies have employed theoretical and experimental approaches to investigate the influence of surface charges upon transmembrane potentials (1,4). However, whether implicitly or explicitly, such studies have assumed that intracellular ion concentrations are unaffected by extracellular fixed charge. However, transmembrane potentials influence transmembrane ion fluxes, and it is therefore necessary to consider whether there are circumstances under which fixed charges may influence intracellular ionic homeostasis.

It is conceivable that the influence of surface charges upon intracellular ionic homeostasis has not previously been investigated because, as shown in Methods, the expectation for a passive ionic distribution between any number of connected compartments is that the equilibrium ion concentrations within any single compartment are uniquely defined with reference to any other single compartment. In other words, each compartment must be at electrochemical equilibrium with each other compartment, irrespective of the number of intervening compartments and the equilibrium ion concentrations within them, as expected from the principle of microscopic reversibility. Thus, if ion concentrations in the extracellular space are fixed, the volume and intracellular ion concentrations of a passive GD cell are uniquely defined by the extracellular environment, irrespective of the ion concentrations within an intervening volume containing fixed charges.

However, ions are not distributed in a purely passive manner in a real cell, and it is therefore necessary to consider the consequences of a charged glycocalyx upon intracellular ionic homeostasis in a more physiologically accurate model cell. This required a method for the calculation of ion concentrations at the outer membrane surface. An immediate theoretical difficulty is that classical GC theory predicts total ion concentrations at the membrane surface that are significantly greater than those in the bulk extracellular space. However, the distribution of charges within a three-dimensional glycocalyx volume raises the possibility that any such changes in local total ion concentration might influence osmotic water movements and therefore glycocalyx volume.

The relationships between glycocalyx fixed charge, volume, and electrical potential

The theoretical approach presented here was developed because the glycocalyx is thick enough in skeletal muscle (some 40–80 nm) to require consideration as a three-dimensional volume rather than as a two-dimensional layer. The GD approach derived in Methods thus incorporates the concept of an osmotically determined glycocalyx volume. With this approach, the glycocalyx charge density (\( \Sigma_g \)) and potential (\( E_2 \)) are dependent variables influenced by the charge valency/osmole of the glycocalyx (\( z_{X(g)} \)). Fig. 2 demonstrates this predicted relationship between \( E_2 \) and \( z_{X(g)} \) for a compartment with a stable osmotic content of molecules (\( X_g \)) of mean charge valency/osmole \( z_{X(g)} \), but which has a volume determined by osmotically driven water movements and thus a variable concentration of \( X_g \) at steady state. The influence of osmotic water movements may be seen by comparison of the predictions of this GD-based approach with those of classical GC theory. To permit comparison of the relationship between the three-dimensional GD approach derived here and two-dimensional GC theory, Appendix 1 shows a derivation of the GC equation using conservation principles similar to those used to derive the GD-based equations. It shows that the novel GD equations reduce to a classical GC function in the limit of an adsorptive layer of zero thickness. Thus, Fig. 3 plots \( E_2 \) against \( \Sigma_g \) from both the GC equation and the GD approach derived here, demonstrating an identical predicted relationship between \( \Sigma_g \) and \( E_2 \) by each approach. However, because \( \Sigma_g = F z_{X(g)} [X_g] \) and \( [X_g] \) decreases as the magnitude of \( z_{X(g)} \) increases (Eq. 12), \( \Sigma_g \) is constrained to a limited range of values at steady state in GD theory. Thus, although both approaches demonstrate a similar relationship between \( \Sigma_g \)
and $E_2$. GD theory additionally suggests a limitation on the physically possible values of $\Sigma_g$. This limitation derives from the alterations in glycocalyceal volume predicted by the GD approach after changes in the value of $z_{X(g)}$.

From a theoretical standpoint (Eq. 12), the two variables that would be expected to influence $[X]_g$, and consequently glycocalyceal volume at constant extracellular osmolarity, are $z_{X(g)}$ and $\eta$ (the mean charge valency of $X_g$ per osmole and the permitted ratio between total ion concentrations in the glycocalyx relative to the bulk extracellular solutions, respectively). Thus, Fig. 4 demonstrates the influences of $z_{X(g)}$ and $\eta$ upon $[X]_g$ (Fig. 4 A) and the resultant value of $[\Sigma]_g$ (Fig. 4 B), the latter being the product $Fz_{X(g)}[X]_g$. This demonstrates that higher magnitudes of $z_{X(g)}$ result in reduced steady-state $[X]_g$ (Fig. 4 A), hence constraining $[\Sigma]_g$ within limits primarily determined by $\eta$ (Fig. 4 B). Since, by definition, the glycocalyx contains a fixed quantity of $X_g$, such reductions in steady-state $[X]_g$ predict that increases in the magnitude of $z_{X(g)}$ would increase glycocalyceal volume from a minimum volume when $z_{X(g)} = 0$.

Electron micrographic analyses of glycocalyceal thicknesses after alterations in extracellular $\text{Ca}^{2+}$ were employed to test this hypothesis that glycocalyceal charge and volume are related. Fig. 5 demonstrates the influence of extracellular $\text{Ca}^{2+}$ upon glycocalyceal volume. Pairs of cutaneous pectoris muscles obtained from different frogs were exposed to each condition of $[\text{Ca}^{2+}]_e$. This demonstrated that there was no significant variation in glycocalyceal thickness, or in the relationship of this thickness to $[\text{Ca}^{2+}]_e$, between muscle fibers from different frogs ($p > 0.05$ in each case). In contrast, pairs of cutaneous pectoris muscles obtained from the same frogs were exposed to different conditions of $[\text{Ca}^{2+}]_e$ and demonstrated increased glycocalyceal thicknesses after both increases (to 5 mM) and decreases (to nominally zero) in $[\text{Ca}^{2+}]_e$.

The biphasic relationship between $[\text{Ca}^{2+}]_e$ and glycocalyceal thickness observed experimentally parallels the relationship between $z_{X(g)}$ and $[X]_g$ shown in Fig. 4. Thus, a possible hypothesis for the observed increases in glycocalyceal diameter in response to both increases and decreases in $[\text{Ca}^{2+}]_e$ from its normal value may be that both maneuvers increase the magnitude of $z_{X(g)}$. In other words, the results suggest that $z_{X(g)}$ may become more negative in low $[\text{Ca}^{2+}]$ and more positive in high $[\text{Ca}^{2+}]$ from an initially intermediate value, and that this is reflected in the observed volume changes. The correspondence between the experimentally observed influence of $[\text{Ca}^{2+}]_e$ on glycocalyceal volume and
The theoretical predictions permits the use of the GD analysis in the further computational studies.

The influence of glyocalyceal potential upon intracellular ion concentrations and \( E_m \)

The CD model of Fraser and Huang (5) was modified as described in Methods to include the glyocalyx GD formulation, thereby allowing simulation of the influence of glyocalyceal charge upon intracellular ion concentrations. Thus, transmembrane ion fluxes were calculated in the CD model from the electrochemical gradients between the bulk intracellular space and the glyocalyx, while the potential and ion concentrations within the glyocalyx were calculated using the GD analysis. Thus, Fig. 6 demonstrates CD modeling of the influence of glyocalyceal charge upon a model resting cell with ion channel and pump density parameters within the physiological range for skeletal muscle. It shows that under physiological conditions, the model predicts that \( z_{X(g)} \) influences \( E_1 \) and \( E_2 \), but not \( E_m \) (Fig. 1).

Fig. 7, A–D, shows an extension of the theoretical analysis to explore the influence of \( z_{X(g)} \) upon critical cellular parameters \( E_1 \), \( E_m \), \( \text{Na}^+/\text{K}^+ \)-ATPase activity (J\(_{\text{Pump}}\)), and intracellular [\( \text{Na}^+ \)] and [\( \text{K}^+ \)], respectively, for model cells with a range of values for the \( \text{Na}^+/\text{K}^+ \)-ATPase density (\( N \)). It shows that in all cases, values of \( N \) above a certain threshold have very little influence on any modeled parameter. This is because the magnitudes of \( E_m \) and the transmembrane ion gradients are then close to their respective maxima and are therefore determined by the thermodynamic constraints upon

\[ \text{Na}^+ / \text{K}^+ \]-ATPase activity (10) rather than by \( N \). When \( N \) is high, more negative values of \( z_{X(g)} \) both increase the steady-state \( \text{Na}^+ / \text{K}^+ \)-ATPase activity and depolarize \( E_1 \), whereas more positive values have the opposite influence. However, despite this significant influence upon \( E_1 \), \( z_{X(g)} \) at higher values of \( N \) has very little influence on \( E_m \). Very low values of \( N \) are not compatible with cell stability. It is only when \( N \) is at an intermediate value—adequate for cellular stability while permitting little increase in pump activity—that \( z_{X(g)} \) may significantly influence \( E_m \).

Within this intermediate range of values of \( N \), the influence of \( z_{X(g)} \) upon \( E_m \) is increased significantly. This is reflected by a relative depolarization of both \( E_1 \) and \( E_m \) at more negative values of \( z_{X(g)} \). The suggestion from Fig. 7 that \( z_{X(g)} \) has its greatest influence upon \( E_m \) under conditions where \( \text{Na}^+ / \text{K}^+ \)-ATPase activity is adequate for cellular stability despite lacking excess capacity invites the question of whether the value of \( N \) is within this intermediate range physiologically. This is unlikely to be the case under normal conditions in skeletal muscle, which is known to have a significant excess \( \text{Na}^+ / \text{K}^+ \)-ATPase capacity (16). However, conditions that reduce \( \text{Na}^+ / \text{K}^+ \)-ATPase activity or increase demand for \( \text{Na}^+ / \text{K}^+ \)-ATPase activity might nevertheless push the effective value of \( N \) into the range within which \( z_{X(g)} \) has a significant influence upon \( E_m \).

Fig. 8 demonstrates one such manipulation of extracellular conditions. It shows the influence of changes in extracellular [\( \text{K}^+ \)] upon \( E_m \). This is a maneuver expected to decrease the energetic favorability of \( \text{Na}^+ / \text{K}^+ \)-ATPase activity, while also increasing passive transmembrane leak currents. Fig. 8 A demonstrates the results of charge-difference modeling of the influence of \( z_{X(g)} \) upon \( E_m \) at different values of [\( \text{K}^+ \)]. from
5 mM to 0.75 mM for a model cell with a constant value of N within the physiological range for skeletal muscle. At higher values of \([K^+]_e\) of 5 mM and 2.5 mM (Fig. 8 A, crosses and squares, respectively), \(z_{X(g)}\) has little influence upon \(E_m\), although, as expected, the lower value of \([K^+]_e\) produces a more polarized model cell. This lack of influence of \(z_{X(g)}\) upon \(E_m\) reflects the fact that \(Na^+/K^+\)-ATPase activity is able to increase at more negative values of \(z_{X(g)}\), thus largely offsetting the resultant increase in leak currents. However, further reduction in \([K^+]_e\) to 1.5 mM or 1 mM permitted two further observations. First, relative depolarization of \(E_m\) was seen with reductions in \([K^+]_e\) at certain (with 1.5 mM \([K^+]_e\)) or all (with 1 mM \([K^+]_e\)) values of \(z_{X(g)}\). This results primarily from a reduction in \(Na^+/K^+\)-ATPase activity at low values of \([K^+]_e\). Second, this reduction in \(Na^+/K^+\)-ATPase activity had the effect of pushing N into the window of values within which \(z_{X(g)}\) was able to have a significant influence on \(E_m\), such that more negative values of \(z_{X(g)}\) produced depolarization. In contrast, this effect was not seen in solutions of higher \([K^+]_e\). Finally, when \([K^+]_e\) was reduced further, the influence of \(z_{X(g)}\) upon \(E_m\) was also reduced, reflecting severe limitation of \(Na^+/K^+\)-ATPase activity at all values of \(z_{X(g)}\).

Fig. 8 B compares predictions from CD modeling with the influence of experimental manipulations of extracellular \([K^+]_e\) and \([Ca^{2+}]_e\). The influence of \([K^+]_e\) upon mean values of \(E_m\) measured between 150 and 300 min after transfer from control to experimental solutions was explored in solutions of normal (1.8 mM) or low (nominally zero) \([Ca^{2+}]_e\). These values were compared to CD modeling of similar manipulations of \([K^+]_e\) in model cells with \(z_{X(g)} = -10\) to mimic the low \([Ca^{2+}]_e\) case, and \(z_{X(g)} = +10\) to mimic the 1.8 mM \([Ca^{2+}]_e\) case. These values are necessarily semiarbitrary because the values of several other key parameters are not known precisely (particularly \(\eta\) and \(N\)). Nevertheless, the use of relatively arbitrary parameters is justified for two reasons. First, the findings are of qualitative value, suggesting that reductions in \([Ca^{2+}]_e\) and more negative values of \(z_{X(g)}\) have similar effects on \(E_m\), as expected. Second, as shown by Figs. 4, 6, and 7 in particular, although the magnitude of the influence of \(z_{X(g)}\) upon \(E_m\) is influenced by the values of \(\eta\) and \(N\), the direction of this influence is not. Therefore, it is possible to conclude that, for example, were \(\eta\) higher than the value of 1.03 used in Fig. 8, smaller changes in \(z_{X(g)}\) would be necessary to achieve the same influence upon \(E_m\). In contrast, there is no way in which, for example, a more negative value

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that model predictions (maximum of $K^+$ predictions. $E_m$ cannot demonstrate precise quantitative agreement between theory and experiment, it does provide qualitative support for the prediction that charge magnitude upon the concentration of mobile ions $E_m$ at intermediate $K^+$m values were stable over extended periods of time (up to 5 h) and a potential difference across the membrane itself ($E_1$) and a potential difference between a region containing fixed charges on or near the outer membrane surface and the bulk extracellular space ($E_2$). In such a scheme, it is the $E_1$ component that constitutes the transmembrane potential actually sensed by integral membrane proteins such as voltage-gated ion channels (3). Previous studies have shown that alterations in extracellular $Ca^{2+}$ reciprocally influence both the voltages, $E_1$ and $E_2$, without altering $E_m$, at least in the short term (1,3,4). However, over longer periods of time, such changes might be expected to influence transmembrane ion fluxes, intracellular ion concentrations, and, hence, $E_m$. This study therefore investigated the influence of extracellular fixed charges upon intracellular ionic homeostasis. It used a novel computational model to explore the effect of alterations in extracellular fixed charge upon intracellular ion concentrations and, hence, $E_m$. The predictions of the model were corroborated by electrophysiological and morphological cellular methods.

A mathematical approach was developed that permitted analytic calculation of the influence of near-surface charges fixed in a layer of nonzero thickness, such as the glyocalyx, upon $E_2$ and local ion concentrations. This theoretical analysis predicted a dependence of glyocalyceal volume upon extracellular $[Ca^{2+}]$, secondary to the influence of fixed charge magnitude upon the concentration of mobile ions within the glyocalyx. Electron micrograph studies showed that glyocalyceal volume was influenced by alterations in extracellular $[Ca^{2+}]$, supporting this theoretical prediction.

The influence of glyocalyceal charge adsorption upon intracellular ionic homeostasis was then explored. The mathematical treatment of the glyocalyx was combined with the CD cellular model of Fraser and Huang (5,7,8), yielding a model that was used to predict the influence of alterations in glyocalyceal charge upon steady-state electrophysiological variables under a variety of conditions. The detailed analysis predicted that although glyocalyceal charge does not influence intracellular ionic homeostasis under normal physiological conditions, thereby leaving $E_m$ unchanged, similar charge effects do significantly influence $E_m$ under conditions where the $Na^+/K^+$-ATPase is moderately challenged.

Experimental measurements of $E_m$ in frog skeletal muscle followed these theoretical predictions closely. Thus, $E_m$ values were stable over extended periods of time (up to 5 h) despite reductions in extracellular $[Ca^{2+}]$ sufficient to produce a more negative glyocalyceal charge. However, the CD modeling suggested that such stability of $E_m$ would have required an increase in $Na^+/K^+$-ATPase activity. This is because more negative glyocalyceal charge causes an increase in near-membrane $[Na^+]$ and $K^+$ that produces a small but significant increase in the transmembrane leak currents of each of these ions according to the Goldman current equation (17,18). CD modeling demonstrated that an

**DISCUSSION**

$E_m$, the electrical potential recorded using intracellular microelectrodes with reference to the bulk extracellular potential, comprises two distinct components: a potential difference across the membrane itself ($E_1$) and a potential difference between a region containing fixed charges on or near the outer membrane surface and the bulk extracellular space ($E_2$). In such a scheme, it is the $E_1$ component that constitutes the transmembrane potential actually sensed by integral membrane proteins such as voltage-gated ion channels (3). Previous studies have shown that alterations in extracellular $Ca^{2+}$ reciprocally influence both the voltages, $E_1$ and $E_2$, without altering $E_m$, at least in the short term (1,3,4). However, over longer periods of time, such changes might be expected to influence transmembrane ion fluxes, intracellular ion concentrations, and, hence, $E_m$. This study therefore investigated the influence of extracellular fixed charges upon intracellular ionic homeostasis. It used a novel computational model to explore the effect of alterations in extracellular fixed charge upon intracellular ion concentrations and, hence, $E_m$. The predictions of the model were corroborated by electrophysiological and morphological cellular methods.
increase in Na\(^+\)/K\(^+\)-ATPase activity, almost sufficient to compensate for the increased leak currents, would be expected to occur due to the influence of the depolarization of \(E_1\) and the fundamental properties of the Na\(^+\)/K\(^+\)-ATPase in model cells with physiological membrane Na\(^+\)/K\(^+\)-ATPase density (10). However, in model cells with reduced Na\(^+\)/K\(^+\)-ATPase density, Na\(^+\)/K\(^+\)-ATPase activity was not able to increase sufficiently to counteract the increased leak fluxes. Consequently, these cells showed a net loss of \([K^+]_i\), a net increase in \([Na^+]_i\), and depolarization of \(E_m\).

Similarly, CD modeling showed that a reduction in extracellular \([K^+]\) could reduce Na\(^+\)/K\(^+\)-ATPase activity sufficiently to prevent an adequate compensatory increase in Na\(^+\)/K\(^+\)-ATPase activity from occurring in model cells with more negative glycocalycal charge. Thus, model cells exposed to both a decrease in \([K^+]_e\) and a more negative glycocalycal charge demonstrated a diminution of \([Na^+]_i\) and \([K^+]_i\) gradients and a depolarization of \(E_m\) with respect to control cells, whereas neither of these maneuvers individually caused depolarization of \(E_m\). This effect was observed at \([K^+]_e\) levels of around 1 mM: lower levels caused depolarization irrespective of glycocalycal charge due to near-complete Na\(^+\)/K\(^+\)-ATPase suppression; higher levels allowed Na\(^+\)/K\(^+\)-ATPase activity to increase in response to depolarization of \(E_1\).

Such theoretical predictions were fulfilled during experimental exposures of frog skeletal muscle fibers to a range of \([K^+]_e\) in the presence of normal and reduced \([Ca^{2+}]_e\). Thus, reduction in extracellular \([Ca^{2+}]\) produced depolarization of \(E_m\) over time only in muscle fibers also exposed to intermediate concentrations of \(K^+\). The experimental results thus supported the finding of the model that limitation of Na\(^+\)/K\(^+\)-ATPase activity by exposure to low \([K^+]_e\) leaves insufficient reserve Na\(^+\)/K\(^+\)-ATPase activity to compensate for the increased transmembrane leak currents provoked by a more negative glycocalycal charge in solutions of reduced \([Ca^{2+}]_e\). This finding that glycocalycal charge can influence intracellular ion concentrations contrasts with assumptions made in the existing literature (4). It suggests, for the first time, that more negative extracellular fixed charge increases Na\(^+\)/K\(^+\)-ATPase activity and thereby ATP usage.

This study thus demonstrates that under normal physiological conditions, extracellular fixed charges do not significantly influence intracellular ion concentrations or \(E_m\), despite their known influence upon the true transmembrane electrical potential (\(E_1\)) (4). However, it also shows that more negative extracellular fixed charges tend to increase passive ion leak currents and Na\(^+\)/K\(^+\)-ATPase activity. This would be expected to increase the metabolic demand upon the cell, or result in cellular K\(^+\) loss and hence depolarization of \(E_m\) if Na\(^+\)/K\(^+\)-ATPase activity was initially near-maximal. These novel findings have possible implications for the influence of \([Ca^{2+}]_e\) upon cells with reduced sodium pump reserve, such as in ischemic tissues, in cardiac failure (12), and under conditions of reduced extracellular \([K^+]\).

**APPENDIX 1: DERIVATION OF THE RELATIONSHIP BETWEEN THE GOUY-CHAPMAN EQUATION AND THE GIBBS-DONNAN ANALYSIS**

The GD analysis considers fixed charges distributed within a three-dimensional volume of space. In contrast, the classical GC equation treats surface charges as being distributed within a two-dimensional layer. It is instructive to compare the predictions of the two approaches, not least because a large body of experimental evidence is in agreement with GC equation under appropriate conditions (9).

To compare such a two-dimensional approach with the three-dimensional approaches considered above, it is helpful to retrieve the GC equation using the same assumptions that were employed above to calculate \(E_2\) by the GD analysis. Thus, the electroneutrality constraint is given by

\[
\sigma = F \int_0^\infty ([A]_x - [B]_x) dx,
\]

where \(\sigma\) represents surface charge density over a two-dimensional surface (C m\(^{-2}\)) and \(x\) denotes distance from the membrane such that the membrane surface is at \(x = 0\). \(\Psi_s\) is thus the potential at a distance \(x\) from the charged surface, such that \(\Psi_s = 0\) is equivalent to \(E_2\). At equilibrium, combining this electroneutrality condition with the requirement of zero net fluxes gives

\[
\Psi_s = -F[C_0] \int_0^\infty \left( e^{\frac{-x^2}{\lambda_D^2}} - e^{\frac{\Psi_s}{F}} \right) dx.
\]

\(\Psi_s\) may then be obtained from the Boltzmann distribution and the Poisson-Boltzmann equation (2):

\[
\Psi_s = \frac{2RT}{F} \ln \left( 1 + \gamma e^{-\frac{x}{\lambda_D}} \right),
\]

where \(\gamma = \tan \left( \frac{\psi_F}{4RT} \right) \) and the Debye length, \(\lambda_D = \sqrt{\frac{\psi_FRT}{2F[C_0]}}\). Note that the Debye length is often taken to be the thickness of the diffuse double layer (2). As shown in full in the purely mathematical argument in Appendix 2, evaluation of the integral gives the GC equation (9):

\[
\Psi_s = \frac{2RT}{F} \arcsinh \left( \frac{\sigma}{\sqrt{8\psi_FRT[C_0]}} \right).
\]

Comparison of the GC equation and Eq. 13 reveals several features of the relationship between potentials generated by two- and three-dimensional charge distributions, respectively. Such a comparison is facilitated if the GC equation is restated in terms of \(\lambda_D\):

\[
\Psi_s = \frac{2RT}{F} \arcsinh \left( \frac{\sigma}{2F[C_0] \lambda_D} \right) = \frac{RT}{F} \arcsinh \left( \frac{\Sigma_m}{2F[C_0] \lambda_D} \right).
\]
Thus, a charge density expressed in two dimensions (C/m²) may be divided by double the Debye length to give an equivalent charge in three dimensions. The latter then produces double the surface potential as would the same concentration of charge in a thick layer. The division of the charge density by double the Debye length is essentially equivalent to integrating the local potential, Ψ, resulting from the charge density over distance, x, from the membrane. The resultant potential is twice that produced by an equivalent density in a layer of thickness significantly greater than the Debye length, reflecting the influence of the low-permittivity membrane in Gouy-Chapman theory. Thus, according to Debye-Hückel theory (2), the potential produced by a given membrane charge at a distance, r, from the membrane, is approximately twice that produced at a similar distance from a similar charge in the bulk extracellular space. This is because ε_m/ε_0 (the permittivity of the membrane relative to that of the extracellular solution) is small.

The above equation may be solved for either σ or Σ to compare the three-dimensional GD approach with the two-dimensional GC approach:

\[
\Sigma_{\theta} = \left(\frac{\sigma}{\lambda_D}\right) \sqrt{1 + \left(\frac{\sigma}{4F[C]\lambda_D}\right)^2}
\]

or

\[
\sigma = \sqrt{8F[C]\lambda_D} \left[\left(1 + \sqrt{1 + \left(\frac{\Sigma_{\theta}}{2F[C]\lambda_D}\right)^2}\right)\right].
\]

These equivalencies were used in Fig. 3 to express σ in terms of the equivalent three-dimensional charge concentration to demonstrate the relationship between E₁ and Σ₁ using the GD and the GC approaches.

**APPENDIX 2: EVALUATION OF THE SURFACE CHARGE INTEGRAL**

For simplicity, the inverse Debye length, κ = 1/λ_D, is used in the derivation

\[
\sigma = -F[C] \int_0^\infty e^{\left(\frac{-\Psi}{RT}\right)} e^{\left(\frac{\Psi}{RT}\right)} dx.
\]

Given \(\frac{\Psi \cdot F}{RT} = 2\ln\left(1 + \gamma e^{-\kappa x}\right)\),

\[
\sigma = -F[C] \int_0^\infty \left(e^{\left(-2\ln\left(1 + \gamma e^{-\kappa x}\right)\right)} - e^{\left(2\ln\left(1 + \gamma e^{-\kappa x}\right)\right)}\right) dx.
\]

Expressing as partial fractions:

\[
\sigma = F[C] \frac{1}{\kappa} \int_1^{1+\gamma} \left(\frac{A}{m} + \frac{B}{m} + \frac{C}{m - 1}\right) \left(\frac{-D}{(2-m)^2} + \frac{E}{m-1} + \frac{G}{(2-m)^2}\right) dm.
\]

Evaluating the partial fraction coefficients:

**Term 1:**

\[
A(m-1) + Bm(m-1) + Cm^2 = m^2 - 4m + 4.
\]

Comparing coefficients in:

\[
m^2 : B + C = 1
\]

\[
m : A - B = -4
\]

const : \(-A - B = 4\)

Evaluating the partial fraction coefficients:

**Term 2:**

\[
\frac{D}{m-1} + \frac{E}{m-1} + \frac{G}{(2-m)^2} = \frac{m^2}{(2-m)^2(m-1)}
\]

\[
D(m-1) + E(2-m)^2 + G(m-1)(2-m) = m^2.
\]

Comparing coefficients in:

\[
m^2 : E - G = 1
\]

\[
m : D - 4E + 3G = 0
\]

const : \(-D + 4E - 2G = 0\)

Substituting back:

\[
\sigma = F[C] \frac{1}{\kappa} \left[\left(\frac{-4}{m} + \frac{1}{m-1}\right) - \left(\frac{4}{(2-m)^2} + \frac{1}{m-1}\right)\right] dm
\]

\[
= F[C] \frac{1}{\kappa} \left[\frac{4(m-2) + 4m^{1+\gamma}}{m^2 - 2m}\right]_1 \]

\[
= F[C] \frac{1}{\kappa} \left[\frac{8\gamma e^{-\kappa \gamma}}{e^{\kappa \gamma} - \gamma}\right]_0
\]

Since \(\gamma = \tanh(\Psi_0 F/4RT)\), and letting \(\xi = \Psi_0 F/4RT\), evaluation of the integral gives:

\[
\sigma = [0] - F[C] \lambda_D \left(\frac{8\tanh\xi}{1 - \tanh^{2}\xi}\right) = F[C] \lambda_D \left(-\frac{8\tanh\xi}{\sech^2\xi}\right)
\]

\[
= F[C] \lambda_D (4\sinh(2\xi)),
\]

\[
\psi_0 = \frac{2RT}{F \arcsinh\left(\frac{\sigma}{\sqrt{8\varepsilon_0\varepsilon_0 RT C_\infty}}\right)}.
\]
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